This Page Is Inserted by IFW Operations and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

As rescanning documents will not correct images, please do not report the images to the Image Problem Mailbox.



(1) Publication number: 0 589 840 A1

12

EUROPEAN PATENT APPLICATION

(21) Application number: 93810653.1

(22) Date of fling: 15.09.93

(i) Int. Cl.⁶: **C12N 15/13,** C12P 21/08, C12N 5/10, C12N 15/62, A61K 39/395, G01N 33/577, G01N 33/68

Declaration under Rule 28(4) EPC (expert solution)

- 30 Priority: 24.09.92 GB 9220228 25.09.92 US 952802
- (3) Date of publication of application: 30.03.94 Bulletin 94/13
- Designated Contracting States:
 AT BE CH DE DK ES FR GB GR IE IT LI LU NL
 PT SE
- (7) Applicant: CIBA-GEIGY AG Klybeckstrasse 141 CH-4002 Basel (CH)
- (1) Applicant: TANOX BIOSYSTEMS, INC. 10301 Stella Link Rd. Houston, TX 77025 (US)

- (72) Inventor: Hardman, Norman Gstaltenrainweg 67 CH-4125 Riehen (CH) Inventor: Kolbinger, Frank Malteserordensstrasse 1D D-79114 Freiburg (DE) Inventor: Saldanha, José 22a Lincoln Way Enfield, Middlesex EN1 1TE (GB)
- (4) Representative: Schluep, Hans-Peter et al c/o CIBA GEIGY AG Patentabteilung Postfach CH-4002 Basel (CH)

- (54) Reshaped monoclonal antibodies against an immunoglobulin isotype.
- (5) The invention relates to reshaped human monoclonal antibodies directed against isotypic determinants of immunoglobulin E (IgE), direct equivalents and derivatives of said antibodies. The molecules of the invention are useful for diagnostics, prophylaxis and treatment of allergy.

The invention relates to reshaped human monoclonal antibodies directed against isotypic determinants of immunoglobulin E (IgE), and derivatives of said antibodies. The antibodies and their derivatives are useful for in vitro and in vivo diagnostics, prophylaxis and treatment of allergy.

Allergy is a hypersensitive state induced by an exaggerated immune response to a foreign agent (the allergen). Immediate (type I) hypersensitivity, characterized by allergic reactions immediately following contact with the allergen, is mediated via B cells and is based on antigen-antibody reactions, whereas delayed hypersensitivity is mediated via T cells and based on mechanisms of cellular immunity. In recent years, the term "allergy" has become more and more synonymous with type I hypersensitivity.

Immediate hypersensitivity is based on the production of antibodies of the immunoglobulin class E (IgE antibodies) by B cells which upon confrontation with the allergen differentiate into antibody secreting plasma cells. The IgE induced reaction is a local event occurring at the site of the allergen's entry into the body, i.e. at mucosal surfaces and/or at local lymph nodes. Locally produced IgE will first sensitize local mast cells, i.e. IgE antibodies bind with their constant regions to Fc, receptors on the surface of the mast cells, and then "spill-over" IgE enters the circulation and binds to receptors on both circulating basophils and tissue-fixed mast cells throughout the body. When the bound IgE is subsequently contacted with the allergen, the Fc, receptors are crosslinked by binding of the allergen whereupon the cells degranulate and release a number of anaphylactic mediators such as histamine, prostaglandins, leukotrienes etc. It is the release of these substances which is responsible for the clinical symptoms typical of immediate hypersensitivity, namely contraction of smooth muscle in the respiratory tract or the intestine, the dilation of small blood vessels and the increase in their permeability to water and plasma proteins, the secretion of mucus resulting e.g. in rhinitis, atopic excema and asthma, and the stimulation of nerve endings in the skin resulting in itching and pain.

In addition, the reaction upon second contact with the allergen is intensified because some B cells form a "memory pool" of surface IgE positive B cells (sigE* B cells) after the first contact with the allergen by expressing IgE on the cell surface.

A promising concept for the treatment of allergy involves the application of monoclonal antibodies, which are IgE isotype-specific and are thus capable of binding IgE. This approach is based on the inhibition of allergic reactions by downregulating the IgE immune response, which is the earliest event in the induction of allergy and provides for the maintenance of the allergic state. As the response of other antibody classes is not affected, both an immediate and a long lasting effect on allergic symptoms is achieved. In addition, antibodies suitable as anti-allergic agents should react with surface IgE positive B cells which into IgE producing plasma cells, so that they can be used to functionally eliminate those B cells. However, antibodies to IgE in principle may also induce mediator release from IgE sensitized mast cells by crosslinking the Fce receptors, thus antagonizing the beneficial effect exerted on the serum IgE and sIgE* B cell level. In consequence, antibodies applicable in therapy of allergy must not be capable of reacting with IgE bound on sensitized mast cells and basophils, but should retain the capability to recognize sIgE* B cells.

Such IgE isotype-specific antibodies have been described e.g. by Chang et al. (Biotechnology 8, 122-126 (1990)), in PCT Application No. 89/06138 and European Patent Application No. 396505. However, as the disclosed antibodies are not of human origin they are less suitable for application to humans due to their immunogenicity as foreign proteins, their rather long persistence in the circulation, and the conceivable formation of damaging immune complexes. These drawbacks may potentially be reduced by transforming e.g. a rodent anti-IgE monoclonal antibody into a chimeric antibody which combines the variable domains of the rodent antibody with human antibody constant domains. This approach conserves the antigen-binding site of the rodent parent anti-IgE antibody, while conferring the human isotype and effector functions. However, for use in humans such a chimeric antibody may not have sufficient clinical advantages over the original rodent antibody.

The immunogenicity of a chimeric antibody can be further reduced by grafting rodent hypervariable regions, also termed complementarity determining regions (CDRs), into the frameworks of human light and heavy chain variable region domains resulting in reshaped human antibodies. The technique involves the substitution or recombinant grafting of antigen-specific rodent CDR sequences for those existent within "generic" human heavy and light chain variable domains (European Patent Application No. 239 400). It is reasoned that this technique will transfer the critical and major portion of the antigen-binding site to the human antibody.

Natural intact immunoglobulins or antibodies comprise a generally Y-shaped tetrameric molecule having an antigen binding-site at the end of each upper arm. An antigen binding site consists of the variable domain of a heavy chain associated with the variable domain of a light chain. More specifically, the antigen binding site of an antibody is essentially formed by the 3 CDRs of the variable domain of a heavy chain (V_L) and the 3 CDRs of the variable domain of the light chain (V_L) . In both V_L and V_R the CDRs alternate with 4 framework regions (FRs) forming a polypeptide chain of the general formula

FR1-CDR1-FR2-CDR2-FR3-CDR3-FR4 (I),

wherein the polypeptide chain is described as starting at the N-terminal extremity and ending at the C-terminal

extremity. The CDRs of V_H and V_L are also referred to as H1, H2, H3, and L1, L2, L3, respectively. The determination as to what constitutes an FR or a CDR is usually made by comparing the amino acid sequences of a number of antibodies raised in the same species and general rules for identification are known in the art ("Sequences of proteins of immunological interest", Kabat E.A. et al., US department of health and human service, Public health service, National Institute of Health).

Recently it has been found that the contribution made by a light chain variable domain to the energetics of binding is small as compared with that made by the associated heavy chain variable domain, and that isolated heavy chain variable domains have an antigen binding activity on their own. Such molecules are commonly referred to as single domain antibodies (Ward, E.S. et al., Nature 341, 544-546 (1989)).

The CDRs form loops which, within the domains, are connected to a b-sheet framework. The relationship between amino acid sequence and structure of a loop can be described by a canonical structure model (Chothia et al., Nature 342, 887-883 (1989)). According to this model, antibodies have only a few main-chain conformations or 'canonical structures' for each hypervariable region. The conformations are determined by the presence of a few key amino acid residues at specific sites in the CDRs and, for certain loops, in the framework regions. Hypervariable regions that have the same conformations in different immunoglobulins have the same or very similar amino acid residues at these sites.

CDR grafting has been carried out for several rodent monoclonal antibodies yielding reshaped human (or humanized) antibodies with a binding affinity significantly lower than that of the rodent CDR-donor antibody. Recent findings have indicated that in addition to the transfer of CDRs changes within the framework of the human sequence may be necessary to provide satisfactory antigen binding activity in the CDR-grafted product.

Queen et al. (Proc. Natl. Acad. Sci. USA 86, 10029-10033 (1989)) have disclosed that the CDRs from a murine anti-Tac monoclonal antibody can be grafted into a human framework. The human frameworks were chosen to maximize homology with the murine sequence. The authors used a computer model of the murine parent antibody to identify amino acid residues located within the FRs that are close enough to interact with the CDRs or antigen. These residues were mutated to the residue found in the murine sequence. The humanized anti-Tac antibody had an affinity that was only about 1/3 that of the murine anti-Tac antibody and maintenance of the human character of this antibody was problematic.

Surprisingly, it has now been found that it is possible to produce reshaped human antibodies directed against human IgE, having an antigen, i.e. IgE, binding affinity which about equals or even exceeds that of the murine CDR-donor antibody.

Accordingly, it is one object of the present invention to provide a reshaped human monoclonal antibody specific for IgE comprising at least one antigen binding site comprising, in sequence, the hypervariable regions CDR1, CDR2 and CDR3; said CDR1 having the amino acid sequence Met-Tyr-Trp-Leu-Glu, said CDR2 having the amino acid sequence Glu-Ile-Ser-Pro-Gly-Thr-Phe-Thr-Thr-Asn-Tyr-Asn-Glu-Lys-Phe-Lys-Ala, said CDR3 having the sequence Phe-Ser-His-Phe-Ser-Gly-Ser-Asn-Tyr-Asp-Tyr-Phe-Asp-Tyr-Phe-Asp-Tyr, said reshaped human antibody having an antigen binding affinity which at least about equals that of the murine CDR-donor antibody, a direct equivalent or a derivative of said reshaped antibody. In the amino acid sequence depicted in SEQ ID NO:1 CDR1 extends from amino acid 31 to 35, CDR2 extends from amino acid 50 to 66 and CDR3 extends from amino acid 99 to 112. The murine CDR-donor antibody is monoclonal antibody TES-C21.

Preferably, the invention relates to a reshaped human antibody comprising at least one antigen binding site comprising:

45

- a) a first domain comprising, in sequence, the hypervariable regions CDR1, CDR2 and CDR3, said CDR1 having the amino acid sequence Met-Tyr-Trp-Leu-Glu, said CDR2 having the amino acid sequence Glu-lie-Ser-Pro-Gly-Thr-Phe-Thr-Thr-Asn-Tyr-Asn-Glu-Lys-Phe-Lys-Ala, said CDR3 having the amino acid sequence Phe-Ser-His-Phe-Ser-Gly-Ser-Asn-Tyr-Asp-Tyr-Phe-Asp-Tyr-Phe-Asp-Tyr; and
- b) a second domain comprising, in sequence, the hypervariable regions CDR1, CDR2 and CDR3, said CDR1 having the amino acid sequence Arg-Ala-Ser-Gln-Ser-Ile-Gly-Thr-Asn-Ile-His, said CDR2 having the amino acid sequence Tyr-Ala-Ser-Glu-Ser-Ile-Ser, said CDR3 having the amino acid sequence Gln-Gln-Ser-Asp-Ser-Trp-Pro-Thr-Thr,
- said reshaped human antibody having an antigen binding affinity which at least about equals that of the murine CDR-donor antibody, a direct equivalent or a derivative of said reshaped antibody. CDR1, CDR2 and CDR3 of the first domain are the CDRs of the protein the sequence of which is identified in SEQ ID NO.1. CDR1, CDR2 and CDR3 of the second domain extend from amino acid 24 to 34, 50 to 56 and 89 to 87, respectively, in the amino acid sequence depicted in SEQ ID NO.3.

When the antigen binding site comprises both the first and second domains, these may be located on the same polypeptide chain or, preferably, each domain may be on a different chain, the first domain being part of an immunoglobulin heavy chain, or fragm in therof, and the second domain being part of an immunoglobulin

light chain or fragment thereof.

30

35

40

45

50

According to the invention a reshaped human antibody refers to a molecule characterized in that (1) it comprises at least one antigen binding site in which each existing chain comprises a human-like framework and (2) any constant region present is at least substantially homologous to, preferably identical with, a human immunoglobulin constant region. As used herein, a "human-like framework" is a framework consisting in sequence of framework regions FR1, FR2, FR3 and FR4, which comprises at least about 70 or more, preferably at least about 75 or more, amino acids identical with those in a framework of a particular human immunoglobulin sequence. Hence, all parts except possibly the CDRs of a reshaped human antibody are substantially homologous to corresponding parts of one or more native human immunoglobulin sequences. For example, a reshaped human antibody would not encompass a chimeric antibody comprising a murine variable region and a human constant region.

A human-like framework may be identical with a framework of a particular human immunoglobulin, or, preferably may differ from the particular human framework, i.e. a limited number of amino acid residues may be inserted, deleted or replaced by other amino acid residues. Such modifications may be confined to a single FR, i.e. FR1, FR2, FR3 or FR4, or involve two, three or all of the four FRs. For example, a hydrophobic amino acid within the human acceptor framework may be replaced with another amino acid, preferably also a hydrophobic amino acid, e.g. a homologous amino acid, replaced with two amino acids, or deleted. Likewise a hydrophilic amino acid within the human framework may be substituted by another amino acid, two amino acids or deleted, whereby replacing amino acids preferably maintain the hydrogen bond structure of the original framework. Such modifications may be performed on a 'trial and error' basis, i.e. the effect thereof is assessed by comparing the antigen-binding affinity of the created reshaped human antibody with that of murine CDRdonor antibody TES-C21. Assays suitable for determination of the antigen binding affinity are described below.

In particular, a limited number of amino acid residues, preferably 1 to 12 residues, within a chosen human acceptor framework may be replaced with amino acid residues present at corresponding positions in a murine monoclonal antibody (human H murine exchange), particularly murine antibody TES-C21, and/or with amino acid residues present at corresponding positions in a different human antibody (human H human exchange). Preferably, the envisaged substitution of (an) amino acid(s) is based on prior identification of particular framework residues to be regarded as potentially crucial for antigen binding and/or V_L/V_H packing. Such crucial amino acids include framework residues which, because of their special nature and/or location:

- are believed to be in contact with, or located near to, amino acids within the CDRs of the antibody;
- could be involved in critical interactions with the antigen;
- are believed to be involved in maintaining the overall integrity of the paired V_H/V_L structure, directly or indirectly influencing interactions within or between the VH and VL domains.

Methods known to be suitable for identification of so-called crucial amino acids include molecular modeling. For example, molecular models of an antigen binding site may be created and displayed on a computer monitor by using computer programs which are generally available and well known to those skilled in the art.

In particular, design of a reshaped antibody of the invention may comprise the following steps:

- a) Construction of a plausible molecular model for V_L and V_H of murine antibody TES-C21, e.g. based on the amino aicd sequences depicted in SEQ ID NOs. 1 and 3 and the corresponding solved structures of a murine antibody determined to be highly homologous by sequence matching. The solved structures may originate from the same murine antibody or from two different murine antibodies.
- b) Selection of suitable human acceptor frameworks from V_H and V_L of known human immunoglobulin sequences, e.g. sequences obtainable from a publicly available database, such as the KABAT database ("Sequences of proteins of immunological interest", Kabat E.A. et al., US department of health and human service. Public health service, National Institute of Health). Suitable human acceptor frameworks are e.g. frameworks from particular immunoglobulins that are highly homologous, preferably unusually homologous as compared with the remaining sequences in the database, to VH and VL domains of antibody TES-C21, or, most preferably consensus frameworks from many human antibodies which are highly homologous to V_H and V_L domains of antibody TES-C21. The heavy and light chain framework sequences chosen for grafting need not be derived from the same human antibody, but preferably are from different human antibodies.
- c) Construction of a molecular model of V_H and V_L of a reshaped human antibody comprising the CDRs of murine TES-C21 and the FRs from the selected human acceptor framework according to formula I.
- d) Comparison of the molecular models obtainable in steps a) and c).

In a reshaped human antibody of the invention, one, some or all of the identified crucial amino acid residues may be substituted with another amino acid residue, in particular with the residue present at that particular position in antibody TES-C21. Preferably, an "original" amino acid residue within the selected human framework is not replaced if it is part of a postulated canonical structure or important in determining the structure of a hypervariable loop. However, substitution of amino acid residues may not be restricted to crucial amino acids. Preferably, changes affecting non-crucial residues are human-human type changes.

In a reshaped human antibody of the invention the amino acid Cys may be in the oxidized state forming -S-S-bridges.

Examples of a reshaped human antibody provided by the present invention include a single domain antibody, a single chain antibody as well as an intact multi-chain, e.g. a tetrameric, antibody comprising full length heavy and light chains and any fragment thereof, e.g. Fv, F(ab')₂, Fab' and Fab fragments.

A single domain antibody comprises a single antigen binding site comprising a single domain.

A single chain antibody (also termed scFv) essentially consists of the variable domains of a heavy and a light chain. Preferably, these variable domains are covalently linked via a short peptide linker comprising from about 10 to 30, particularly about 15 amino acids selected from glycine and serine. A preferred peptide linker is the 15 amino acid polypeptide consisting of three repetitive units of Gly-Gly-Gly-Gly-Ser. A single chain antibody does not include a constant part of either heavy or light chain.

The reshaped human antibody of the invention is specific for IgE, i.e. it is directed against an isotypic determinant of human IgE. Accordingly, the antibody of the invention recognizes an antigenic determinant on the e heavy chain common to immunoglobulins of class IgE, i.e. it reacts with IgE molecules of different specificities but does not react with immunoglobulins of other isotypes or with immunoglobulin light chains.

A reshaped antibody of the invention is required to have an IgE-binding affinity which at least about equals that of murine antibody TES-C21. As used herein before or hereinafter, the term "at least about equals" means that the IgE-binding affinity of the reshaped human antibody (test antibody), on a statistical basis, is at least about 90 %, preferably higher than 90 %, particularly within about 100 % and about 250 %, of reference antibody TES-C21. A reshaped antibody is to be compared against the corresponding structure of TES-C21. For example, if the reshaped antibody is a single domain antibody its affinity should be related to single domain TES-C21. This murine single domain antibody can be easily prepared based on the information given in SEQ ID NOs. 1 and 3. In the description, no distinction is made between "affinity" or "avidity" of an antibody, but the term "affinity" is to refer to either affinity or avidity.

Determination of affinities of the reference and the reshaped test antibody is to be performed in the same fashion, i.e. under identical conditions in the same assay. The antibodies compared with each other should have about the same degree of purity. It is preferred to use highly purified antibodies.

The binding affinity of an antibody for IgE is determined using a suitable quantitative assay which can be easily established by a person with ordinary skill in the art based on known techniques and principles.

A suitable parameter to be determined is the equilibrium constant $K_{\rm aff}$ (affinity constant). A variety of mathematical equations have been developed to facilitate experimental calculations of affinity constants for the antibody-antigen interaction. Suitable experimental methods for the measurement of $K_{\rm aff}$ may e.g. rely on the measurement of the bound to free antigen ratio, e.g the competitive radioimmunoassay (RIA) or the competitive enzyme-linked immunoadsorbent assay (EIA), or on the measurement of the total antibody concentration, e.g. the non-competitive, solid phase EIA described by Beatty et al. (J. Immunol. Meth. 100, 173-179 (1987)).

Preferably, K_{eff} is determined analyzing real-time biospecific interaction (Jönsson, U. et al., Biotechniques 11, 620-627 (1991) of the antibody with the IgE antigen on a BIA core™ system using CM5 surface chips (Pharmacia Biosensor, Uppsala, Sweden). The assay is essentially performed according to the manufacturer's instruction and involves determination of the kinetic constants k_{sss} and k_{diss}. A suitable antigen is e.g. commercially available human IgE provided e.g. by Serotec (e.g. BP 094, Dottikon Switzerland) or a chimeric antibody having a human e constant region such as SE 44 (Sun et al., J. Cell. Biol. 109, 289a (1989)). In particular, this assay comprises an experimental cycle comprising:

- 1) Immobilization of a so-called catching antibody on the chip surface by chemical means, i.e. for measurements involving murine antibody TES-C21 an anti-mouse antibody, e.g. anti-mouse IgG, or for measurements involving a reshaped human antibody of the invention an anti-human antibody, e.g. anti-human IgG is employed.
- 2) Binding reference or test antibody to the immobilized catching antibody

45

3) Contacting the bound reference or test antibody with a fixed concentration of antigen.

Preferably several, e.g. four, experimental cycles are performed using a constant amount of bound antibody and varying the (known) concentration of IgE. After completion of each cycle the surface is regenerated, e.g. with an acid such as HCl.

Design of a reshaped antibody of the invention aims at constructing an antibody exhibiting a high association rate (k_{ess}), preferably 2.5 x 10⁶ M⁻¹s⁻¹ or higher, combined with a low dissociation rate (k_{ess}), preferably 1.9 x 10⁻⁶s⁻¹ or lower.

As used herein, a direct equivalent of a rehaped human antibody of the invention is a reshaped human antibody comprising, in sequence, CDR1, CDR2 and CDR3 as shown in SEQ ID NO. 1 and, optionally, CDR1, CDR2 and CDR3 as shown in SEQ ID NO. 3, wherein within one variable domain up to four amino acid residues

within the CDRs, i.e. one two, three or four amino within the CDRs are replaced with another amino acid. Thus, by the term "direct equivalents thereof" is meant either a single domain reshaped human antibody (protein Y)

- (1) in which the hypervariable regions CDR1, CDR2 and CDR3 taken as a whole are at least 90 % homologous to the CDRs as shown in SEQ ID NO. 1 and,
- (2) which has an affinity for IgE which at least about equals that of the reference protein of the invention having FRs identical to those of protein Y but having CDRs identical with those in SEQ ID NO.1; or a reshaped antibody having two domains per binding site (protein Y')
 - (1) in which the hypervariable regions CDR1_H, CDR2_H, CDR3_H and CDR1_L, CDR2_L, CDR3_L taken as a whole are at least 80 %, preferably 90 % homologous to the CDRs as shown in SEQ ID NOs. 1 and 3, and (2) which has an affinity for IgE which at least about equals that of the reference protein of the invention having FRs and constant parts identical to those of protein Y but having hypervariable regions CDR1_H, CDR2_H, CDR3_H and CDR1_L, CDR3_L identical with those in SEQ ID NOs.1 and 3.

The latter criterion can be tested by determining K_{eff}, e.g. according to the method described above. Murine monoclonal antibody TES-C21 displays (among others) the following characteristics, which are also common to a reshaped human antibody of the present invention:

- it inhibits the binding of IgE to cells bearing Fc, receptors I or II;
- binds specifically to human-IgE secreting cells;
- does not recognize and bind IgE bound on the surface of cells bearing Fc_e receptors I or II, for example sensitized mast cells and basophils,
- does not trigger mediator (e.g. histamine) release.
- inhibits IgE formation in the immune response.

25

30

35

These characteristic abilities can be determined by methods known in the art, e.g. those disclosed in European Application No. 396505 which is herein incorporated by reference.

A preferred reshaped antibody, or a derivative thereof of the invention comprises at least:

- a) one immunoglobulin heavy chain or a fragment thereof which comprises a variable domain comprising in sequence the hypervariable regions CDR1_H. CDR2_H and CDR3_H and the constant part or fragment thereof of a human heavy chain, said CDR1_H having the amino acid sequence Met-Tyr-Trp-Leu-Glu, said CDR2_H having the amino acid sequence Glu-Ile-Ser-Pto-Gly-Thr-Phe-Thr-ThrAsn-Tyr-Asn-Glu-Lys-Phe-Lys-Ala, said CDR3_H having the sequence Phe-Ser-His-Phe-Ser-Gly-Ser-Asn-Tyr-Asp-Tyr-Phe-Asp-Tyr-Phe-Asp-Tyr-Phe-Asp-Tyr: and
- b) and one immunoglobulin light chain, or a fragment thereof, comprising a light chain variable domain comprising in sequence the hypervariable regions CDR1_L,CDR2_L and CDR3_Land the constant part, or a fragment thereof, of a human light chain, said CDR1_L having the amino acid sequence Arg-Ala-Ser-Gln-Ser-Ile-Gly-Thr-Asn-Ile-His, said CDR2_L having the amino acid sequence Tyr-Ala-Ser-Glu-Ser-Ile-Ser, said CDR3_L having the amino acid sequence Gln-Gln-Ser-Asp-Ser-Trp-Pro-Thr-Thr.

A fragment of an immunoglobulin heavy or light chain is a heavy or light chain which is not a full length chain and comprises a variable domain and optionally part of the constant part of the chain.

More preferred is a reshaped human antibody, or a derivative thereof, comprising at least

- a) one heavy chain comprising a variable domain having an amino acid sequence substantially identical with that shown in SEQ ID NO. 11 starting with the amino acid at position 1 and ending with the amino acid at position 123 and the constant part of a human heavy chain; and
- b) one light chain comprising a variable domain having an amino acid sequence substantially identical with that shown in SEQ ID NO. 5 starting with the amino acid at position 1 and ending with the amino acid at position 107 and the constant part of a human light chain;
- Particularly preferred is a reshaped human antibody or a derivative thereof comprising at least
- a) one heavy chain comprising a variable domain having an amino acid sequence substantially identical with that shown in SEQ ID NO. 13 starting with the amino acid at position 1 and ending with the amino acid at position 123 and the constant part of a human heavy chain; and
- b) one light chain comprising a variable domain having an amino acid sequence substantially identical with that shown in SEQ ID NO. 5 starting with the amino acid at position 1 and ending with the amino acid at position 107 and the constant part of a human light chain;

The residue designations given in the present application correspond with the linear numbering of the ami-

The constant part of a human heavy chain can be selected from any of the isotypes alpha (a), delta (d), gamma(g) or $mu(\mu)$. Heavy chains of various subclasses such as the IgG subclasses 1-4 can be used. Preferred is the constant part of the human gl chain. The different classes and subclasses of heavy chains are involved in different effector functions and thus, by choosing the type of the heavy chain constant region, reshaped human antibodies with the desired effector functions can be produced. The constant part of a light chain

is a lambda(i), or preferably a kappa(k) chain.

45

Most preferred is reshaped human antibody designated H3L1 produced by the cell line EH31.8.

The invention also concerns a derivative of a reshaped human antibody of the invention. A derivative of a reshaped human antibody has the antigenic specificity of said antibody. According to the invention a derivative is meant to be any molecule obtainable by modification of an antibody of the invention, e.g. by adsorption or chemical modification. For example, depending on the intended use of the derivative, an antibody of the invention may be derivatized by covalent or non-covalent attachment of another proteinaceous or non-proteinaceous molecule. Covalent attachment resulting in antibody conjugates is achieved e.g. using coupling techniques known in the art. In such conjugates, the antibody is bound to the conjugation partner directly or by way of a spacer or linker group. Examples of derivatives are radioactively labelled reshaped human antibodies and conjugates of a reshaped human antibody of the invention, e.g. with an enzyme, a fluorescent or chemiliuminescent marker, a suitable cytotoxic or cytostatic substance, a metal chelate, a protein that is not an enzyme such as avidin, or with a non-proteinaceous molecule such as biotin.

Enzymes used for antibody conjugates of the invention are, for example, horseradish peroxidase, alkaline phosphatase, b-D-galactosidase, glucose oxidase, glucoamylase, carbonic anhydrase, acetylcholinesterase, lysozyme, malate dehydrogenase or glucose-6-phosphate dehydrogenase.

Fluorescent markers include fluorescein, fluorochrome, rhodamine, and the like.

Chemiluminescence markers are e.g. acridinium esters of luminol.

Examples of metal chelates are ethylenediaminetetraacetic acid (EDTA), diethylenetriaminepentaacetic acid (DPTA), 1,4,8,11-tetraazatetradecane, 1,4,8,11-tetraazatetradecane-1,4,8,11-tetraacetic acid, 1-oxa-4,7,12,15-tetraazateptadecane-4,7,12,15-tetraacetic acid, or the like.

Radioactively labelled antibodies or fragments of the invention contain e.g. radioactive iodine (1231, 1251, 1311), tritium (3H), carbon (14C), sulfur (35S), yttrium (90Y), technetium (90mTc), or the like.

The invention further concerns a method for the manufacture of anti-IgE reshaped human antibodies, direct equivalents and derivatives thereof according to the invention.

The reshaped human antibody, a direct equivalent or a derivative thereof according to the invention is prepared by a process that is known per se, characterized in that suitable host cells as defined further below producing a protein of the invention, are multiplied in vitro or in vivo and, if required, the desired protein is isolated and, optionally, converted into a derivative thereof. A protein of the invention can be prepared by a process comprising culturing any suitable transformable host under conditions which allow the expression of said protein, isolating said protein and, optionally, converting the isolated protein into another protein of the invention, e.g. by proteolytic cleavage, or into a derivative of the invention, e.g. by attachment of another compound, e.g. a protein or a non-proteinaceous molecule, as mentioned above.

In a preferred embodiment of the invention, there is provided a process for producing a multi-chain anti-IgE reshaped human antibody which comprises (1) culturing a suitable host cell which has been transformed with first and second DNA constructs of the invention as defined below and (2) recovering an active anti-IgE reshaped human antibody from the culture. In this context an active antibody is an antibody specifically binding to IgE. A multi-chain antibody is an antibody comprising at least one antigen-binding site comprising a heavy and a light chain variable domain.

Alternatively, the heavy and light chain may be separately recovered and reconstituted into an active antibody after in vitro folding. Appropriate reconstitution methods are well known in the art. Therefore a process for producing a multi-chain antibody of the invention may also comprise:

- (1) culturing a first host cell which is transformed with a first DNA construct of the invention and recovering said heavy chain or fragment thereof from the culture and
- (2) culturing a second host cell which is transformed with a second DNA construct of the invention and recovering said light chain or fragment thereof from the culture and
- (3) reconstituting in vitro an active anti-IgE reshaped antibody from the heavy chain or fragment thereof obtained in (1) and the light chain or fragment thereof obtained in (2).

In a similar manner, there is also provided a process for producing a single chain or a single domain reshaped human antibody of the invention which comprises (1) culturing a host cell which is transformed with a DNA construct respectively encoding a single chain or single domain reshaped human antibody of the invention and (2) recovering said polypeptide from the culture.

Fragments of the reshaped human antibodies, for example Fab, Fab' or F(ab')₂ fragments, can be prepared by recombinant DNA techniques as described above or from an intact multichain reshaped human antibody prepared as mentioned above by methods known per se, e.g. by digestion with enzymes such as papain or pepsin and/or cleavage of disulfide bonds by chemical reduction.

Suitable host cells include eukaryotic cells, e.g. animal cells, plant cells and fungi, and prokaryotic cells, such as gram-positive and gram-negative bacteria, e.g. E. coli. Preferred eukaryotic host cells are cells of mam-

malian origin and yeast cells.

5

As used hereinbefore or hereinafter, <u>in vitro</u> means <u>ex vivo</u>, thus including e.g. cell culture and tissue culture conditions.

For example, multiplication of mammalian cells in vitro is carried out in suitable culture media, which are the customary standard culture media, such as Dulbecco's Modified Eagle Medium (DMEM) or RPMI 1640 medium, optionally replenished by a mammalian serum, e.g. fetal calf serum, or trace elements and growth sustaining supplements, e.g feeder cells such as normal mouse peritoneal exudate cells, spleen cells, bone marrow macrophages, 2-aminoethanol, insulin, transferrin, low density lipoprotein, oleic acid, or the like.

In vitro production provides relatively pure antibody preparations and allows scale-up to give large amounts of the desired antibodies. Techniques for bacterial cell, yeast and mammalian cell cultivation are known in the art and include homogeneous suspension culture, e.g. in an airlift reactor or in a continuous stirrer reactor, or immobilized or entrapped cell culture, e.g. in hollow fibres, microcapsules, on agarose microbeads or ceramic cartridges.

Large quantities of the desired reshaped human antibodies of the invention can also be obtained by multiplying mammalian cells in vivo, For this purpose, hybridoma cells producing the desired antibodies are injected into histocompatible mammals to cause growth of antibody-producing tumours. Optionally, the animals are primed with a hydrocarbon, especially mineral oils such as pristane (tetramethyl pentadecane), prior to the injection. After one to three weeks, the antibodies are isolated from the body fluids of those mammals. For example, transfected cells derived from hybridoma cell line Sp2/0 that produce the desired antibodies are injected intraperitoneally into Balb/c mice optionally pre-treated with pristane, and, after one to two weeks, ascitic fluid is taken from the animals.

The cell culture supernatants are screened for the desired reshaped human antibodies, preferentially with an enzyme immunoassay, e.g. a sandwich assay or a dot-assay, or a radioimmunoassay using human IgE as antigen. For example, a sandwich enzyme immunoassay may be used to determine whether correctly assembled immunoglobulins are present in cell culture supernatants, whereby an antibody directed to the light chain human constant region k or I (as appropriate) and another antibody directed to the heavy chain human constant region e.g. g of the desired subclass are used, one of which is coated to a solid support and the other one conjugated to an enzyme allowing detection with a suitable enzyme substrate. Such an immunoassay is, for example, an enzyme-linked immunoadsorbent assay (ELISA) wherein a suitable carrier, e.g. plastic microtitre plates are coated with immunoglobulin E and incubated with the culture supernatant to be tested. Bound monoclonal antibodies are detected by incubation with an enzyme-labelled antibody recognizing the anti-IgE antibodies in the supernatant and by subsequent addition of an appropriate enzyme substrate solution. The enzyme substrate reaction results, for example, in a color change which can be observed by eye or with optical measuring devices.

For isolation of the reshaped human antibodies, the immunoglobulins in the culture supernatants or in the ascitic fluid may be concentrated, e.g. by precipitation with ammonium sulphate, dialysis against hygroscopic material such as PEG, filtration through selective membranes, or the like. If necessary and/or desired, the antibodies are purified by the customary chromatography methods, for example gel filtration, ion-exchange chromatography, hydrophobic interaction chromatography or affinity chromatography, e.g. immunoaffinitychromatography. Preferably, the reshaped human antibodies are isolated from cell supernatants containing them by a procedure comprising a chromatographic purification step, e.g. affinity chromatography, for example with Protein A (if the antibody of the invention comprises an Fc part), ion-exchange chromatography, and/or gel filtration.

The reshaped human antibody derivatives of the invention are prepared by methods known per se, e.g. by adsorption of the reshaped human antibodies to another compound or by coupling providing chemically bound conjugates. Conjugates of antibodies of the invention with a protein, e.g. an enzyme are prepared e.g. by reacting an antibody prepared as described above with the protein in the presence of a coupling agent, e.g. glutaraldehyde, periodate, N,N'-o-phenylenedimaleimide, N-(m-maleimidobenzoyloxy)-succinimide, N-(3-[2'-pyridyldithio]-propionoxy)-succinimide, N-ethyl-N'-(3-dimethylaminopropyl)-carbodiimide or the like. Conjugates with biotin are prepared e.g. by reacting antibodies with an activated ester of biotin such as the biotin N-hydroxysuccinimide ester. Conjugates with fluorescent or chemiluminescent markers are prepared in the presence of a coupling agent, e.g. those listed above, or by reaction with an isothiocyanate, preferably fluorescein-isothiocyanate.

Reshaped antibodies radioactively labelled with iodine (1231, 1261, 1311) are obtained from the antibodies of the invention by iodination according to methods known <u>per se</u>, for example with radioactive sodium or potassium iodide and a chemical oxidizing agent, such as sodium hypochlorite, chloramine T or the like, or an enzymatic oxidizing agent, such as lactoperoxidase, or glucose oxidase and glucose. Antibodies or fragments according to the invention are coupled to yttrium (20Y) for example by diethylenetriaminepentaacetic acid

(DPTA)-chelation. Technetium-99m labelled antibodies or fragments are prepared by ligand exchange processes, for example by reducing pertechnate (TcO₄⁻) with stannous ion solution, chelating the reduced technetium onto a Sephadex column and applying the antibodies to this column, or by direct labelling techniques, e.g. by incubating pertechnate, a reducing agent such as SnCl₂, a buffer solution such as sodium-potassium phthalate solution, and the antibodies of the invention.

Conjugates of antibodies of the invention to a protein may also be prepared directly by recombinant DNA techniques, e.g. those described below.

The process for producing a reshaped human antibody, a direct equivalent or a derivative according to the invention should yield the desired protein in an amount sufficient for affinity and specificity determinations.

The invention also concerns recombinant DNAs coding for the reshaped human antibodies of the invention and direct equivalents thereof. In a very general manner, there are provided DNA molecules encoding a single domain reshaped human antibody of the invention, a single chain reshaped human antibody of the invention, a heavy or light chain or fragments thereof of a reshaped human antibody of the invention. By definition such DNAs comprise coding single stranded DNAs, double stranded DNAs consisting of said coding DNAs and complementary DNAs thereto, or these complementary (single stranded) DNAs themselves. More specifically, the invention relates to first and second DNA constructs as described below.

The first DNA construct encodes a heavy chain or a fragment thereof and comprises

20

25

30

35

45

a) a first part which encodes a variable domain comprising alternatively FRs and CDRs, said CDRs being in sequence CDR1_H, CDR2_H and CDR3_H, the amino acid sequences of which in SEQ ID NO.1 extend from positions 31 to 35, 50 to 66 and 99 to 112, respectively; this first part starting with a codon encoding the first amino acid of the variable domain and ending with a codon encoding the last amino acid of the variable domain, and optionally,

b) a second part encoding a human heavy chain constant part or fragment thereof which starts with a codon encoding the first amino acid of the constant part of the heavy chain and ends with a codon encoding the last amino acid of the constant part or fragment thereof, followed by a nonsense codon. Preferably, this first part encodes a variable domain having an amino acid sequence substantially identical to the amino acid sequence depicted in SEQ ID NO. 13 starting with the amino acid at position 1 and ending with the amino acid at position 123. More preferably, the first part has the nucleotide sequence as shown in SEQ ID NO. 13 starting with the nucleotide at position 79 and ending with the nucleotide at position 447. The second part may be a DNA fragment of genomic origin (comprising introns) or a cDNA fragment (without introns). If present, a second part encoding the constant part of the gl chain is preferred.

The second DNA construct encodes a light chain or a fragment thereof and comprises

a) a first part which encodes a variable domain comprising alternatively FRs and CDRs, said CDRs being in sequence CDR1_L, CDR2_L and CDR3_L, the amino acid sequences of which in SEQ ID NO. 3 extend from positions 24 to 34, 50 to 56 and 89 to 97, respectively; this first part starting with a codon encoding the first amino acid of the variable domain and ending with a codon encoding the last amino acid of the variable domain, and optionally

b) a second part encoding a human light chain constant part or fragment thereof which starts with a codon encoding the first amino acid of the constant part of the light chain and ends with a codon encoding the last amino acid of the constant part or fragment thereof, followed by a nonsense codon. Preferably, this first part encodes a variable domain having an amino acid sequence substantially identical to the amino acid sequence depicted in SEQ ID NO. 5 starting with the amino acid at position 1 and ending with the amino acid at position 107. More preferably, the first part has the nucleotide sequence as shown in SEQ ID NO. 5 starting with the nucleotide at position 82 and ending with the nucleotide at position 403. The second part may be a DNA fragment of genomic origin (comprising introns) or a cDNA fragment (without introns). If present, a second part encoding the constant part of the κ chain is preferred.

Preferred are first and second DNA constructs comprising both the first and the second part. In this case the first and second parts may be separated by intron sequences.

Advantageously, the first and second DNA construct comprise a third part which is located upstream of the first part and which encodes a leader peptide; this third part starts with the codon encoding the first amino acid and ends with a codon encoding the last amino acid of the leader peptide. A suitable leader peptide is a peptide required for secretion of the chains by the host organism in which they are expressed and which is subsequently removed by the host. Preferably, the third parts of the first and second DNA constructs encode a leader peptide of an immunoglobulin gene. Most preferably, the third part of the first DNA construct encodes a leader peptide having an amino acid substantially identical with the sequence shown in SEQ ID NO. 13, starting with the amino acid at position -19 and ending with the amino acid at position -1. Also most preferably, the third part of the second DNA construct encodes a leader peptide having an amino acid substantially identical with the sequence shown in SEQ ID No. 5, starting with the amino acid at position -20 and ending with the

amino acid at position -1.

15

30

35

The invention also concerns a recombinant DNA coding for a direct equivalent of a reshaped human antibody of the invention and a recombinant DNA coding for a conjugate of an antibody of the invention to a protein.

The present state of the art is such that a person with ordinary skill in the art will be able to synthesize the DNA molecules of the invention given the written information provided herein, i.e. the amino acid sequences of the CDRs and the DNA sequences coding therefor (SEQ ID NOs. 1 and 3). A suitable method for obtaining a DNA construct encoding a variable domain of a reshaped human antibody of the invention comprises the synthesis of a number of oligonucleotides, their amplification by the PCR method, and their splicing to give the desired DNA sequence. An alternative method for constructing a variable domain gene comprises:

- doning a gene encoding a human monoclonal antibody of whatever specificity,
- determining the DNA segments encoding the FRs and CDRs,
- removing the DNA segments encoding the CDRs, so that the DNA segments encoding the FRs are fused together with suitable restriction sites at the junctions,
- preparing double stranded synthetic CDR cassettes according to the above identified sequences in SEQ ID Nos. 1 and 3, said cassettes having sticky ends,
- ligating the cassettes at the junctions of the FRs (European Patent Application No. 239 400).

If desired, the DNA constructs of the invention may be mutated by a variety of well-known standard procedures, e.g. by inducing random mutations or by site-directed mutagenesis. In a DNA construct coding for a reshaped human antibody of the invention mutagenesis may not lead to an alteration of any amino acid located within a CDR. In a DNA construct coding for a direct equivalent of a reshaped antibody of the invention a replacement of a nucleotide with another nucleotide may alter the amino acid sequence in one or more CDRs.

A DNA coding for a direct equivalents of the reshaped antibodies of the invention may be prepared according to procedures known in the art e.g. by random or site-directed mutation of a DNA coding for a reshaped antibody of the invention. A mutation which is not a silent mutation but results in the replacement of at least one amino acid residue located within a CDR may yield a DNA coding for a direct equivalent of a reshaped antibody of the invention, if the protein thus produced meets the above-mentioned criterion.

As used in the following part of the specification, a reshaped human antibody of the invention is meant to include direct equivalents thereof.

Furthermore the invention concerns a recombinant DNA which is a hybrid vector comprising at least one of the above described DNA constructs, e.g. an insert coding for a light chain variable domain and/or a heavy chain variable domain, said vector being capable of replicating in a prokaryotic and/or eukaryotic host.

Preferred hybrid vectors of the invention comprise an insert coding for a light chain as described hereinbefore, and/or an insert coding for a heavy chain as described hereinbefore.

The hybrid vectors of the invention comprise an origin of replication or an autonomously replicating sequence, one or more dominant marker sequences and, optionally, expression control sequences, signal sequences and additional restriction sites.

Preferably, the hybrid vector of the invention comprises an above-described insert operably linked to an expression control sequence, in particular those described hereinafter.

Vectors typically perform two functions in collaboration with compatible host cells. One function is to facilitate the cloning of the nucleic acid that encodes the immunoglobulin chain, i.e. to produce usable quantities of the nucleic acid (cloning vectors). The other function is to provide for replication and expression of the gene constructs in a suitable host, either by maintenance as an extrachromosomal element or by integration into the host chromosome (expression vectors). A cloning vector comprises the gene constructs as described above, an origin of replication or an autonomously replicating sequence, selectable marker sequences and, optionally, signal sequences and additional restriction sites. An expression vector additionally comprises expression control sequences essential for the transcription and translation of the genes.

An origin of replication or an autonomously replicating sequence is provided either by construction of the vector to include an exogeneous origin such as derived from Simian virus 40 (SV 40) or another viral source, or by the host cell chromosomal mechanisms.

The markers allow for selection of host cells which contain the vector. Selection markers include genes which confer resistance to heavy metals such as copper or to antibiotics such as tetracycline, ampicillin, geneticin (G-418), neomycin, kanamycin or hygromycin, or genes which complement a genetic lesion of the host cell such as the absence of thymidine kinase, hypoxanthine phosphoryl transferase, dihydrofolate reductase or the like

Signal sequences may be, for example, presequences or secretory leaders coding for a leader peptide directing the secretion of the antibody, splice signals, or the like.

As expression control sequences, the vector DNA comprises a promoter, sequences necessary for the

initiation and termination of transcription and for stabilizing the mRNA and, optionally, enhancers and further regulatory sequences. A wide variety of promoting sequences may be employed, depending on the nature of the host cell. Promoters that are strong and at the same time well regulated are the most useful. Sequences for the initiation of translation are for example Shine-Dalgarno sequences. Sequences necessary for the initiation and termination of transcription and for stabilizing the mRNA are commonly available from the noncoding 5'-regions and 3'-regions, respectively, of viral or eukaryotic cDNAs, e.g. from the expression host. Enhancers are transcription-stimulating DNA sequences of genomic or viral origin, e.g. derived from Simian virus, polyoma virus, bovine papilloma virus, Moloney sarcoma virus, or particularly from human cytomegalovirus.

The various DNA segments of the vector DNA are operationally linked, i.e. they are contiguous and placed into a functional relationship with each other.

Examples of vectors which are suitable for replication and expression in an $\underline{E.~coli}$ strain are bacteriophages, for example derivatives of λ bacteriophages, or plasmids.

Sultable vectors comprise a complete replicon, a marker gene, recognition sequences for restriction endonucleases, so that the foreign DNA and, if appropriate, the expression control sequence can be inserted at these sites, and optionally signal sequences and enhancers. An expression vector according to the invention comprises an expression cassette comprising a suitable promoter and a DNA construct as defined above, which DNA is controlled by said promoter.

Microbial promoters are, for example, the strong leftward promoter P_L of bacteriophage λ which is controlled by a temperature sensitive repressor. Also suitable are \underline{E} . \underline{coli} promoters such as the lac (lactose) promoter regulated by the lac repressor and induced by isopropyl- β -D-thiogalactoside, the trp (tryptophan) promoter regulated by the trp repressor and induced e.g. by tryptophan starvation, and the tac (hybrid trp-lac promoter) regulated by the lac repressor.

Vectors which are suitable for replication and expression in yeast contain a yeast replication start and a selective genetic marker for yeast. One group of such vectors includes so-called ars sequences (autonomous replication sequences) as origin of replication. These vectors are retained extrachromosomally within the yeast cell after the transformation and are replicated autonomously. Furthermore, vectors which contain all or part of the 2 μ (2 mikron) plasmid DNA from Saccharomyces cerevisiae can be used. Such vectors will get integrated by recombination into 2 μ plasmids already existing within the cell, or replicate autonomously. 2 μ sequences are particularly suitable when high transformation frequency and high copy numbers are to be achieved.

Expression control sequences which are suitable for expression in yeast are, for example, those of highly expressed yeast genes. Thus, the promoters for the <u>TRP1</u> gene, the <u>ADHI</u> or <u>ADHII</u> gene, acid phosphatase (<u>PHO3</u> or <u>PHO5</u>) gene, isocytochrome gene or a promoter involved with the glycolytic pathway, such as the promoter of the enolase, glyceraldehyde-3-phosphate kinase (<u>PGK</u>), hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triose-phosphate isomerase, phosphoglucose isomerase and glucokinase genes, can be used.

Promoters suitable for mammalian host cells are obtainable from a human immunoglobulin gene or from viruses such as Simian virus 40 (SV 40), Rous sarcoma virus (RSV), adenovirus 2, bovine papilloma virus (BPV), papovavirus BK mutant (BKV), or mouse or human cytomegalovirus (CMV). Preferred is the human CMV promoter. Alternatively, the vectors may comprise promoters from mammalian expression products, such as actin, collagen, myosin etc., or the native promoter and control sequences which are normally associated with the immunoglobulin gene sequences.

The vectors may be suitable for both prokaryotic and eukaryotic hosts. Once a DNA molecule of the invention is prepared it may be conveniently transferred into a suitable expression vector. Expression vectors comprising a suitable promoter or genes encoding heavy or light chain constant parts are publicly available.

The gene constructs for the light chain and for the heavy chain are sequentially or simultaneously transferred into the host cells with the help of two vectors. Alternatively, both heavy and light chains are cloned into the same hybrid vector and incorporated in a one step-procedure as a single construct into the host cells. A third alternative utilises co-transfection of unlinked DNA fragments.

The recombinant DNAs coding for the desired reshaped human antibody can be prepared, for example, by culturing a transformed host cell.

In particular, such DNAs can be prepared by a method comprising

- a) preparing DNA coding for the variable heavy and/or light chain variable domain of a reshaped human antibody specific for IgE
- b) preparing DNA coding for the heavy and/or light chain constant region of a human antibody, e.g. by isolating DNA from a genomic library and selecting the desired DNAs coding for said constant regions of antibodies using DNA probes;
- c) incorporating the DNA of step a) or the DNA of steps a) and b) into appropriate hybrid vectors;
- d) transferring the obtained hybrid vectors into a recipient host cell or retrieving the DNA coding for the

desired genes and transferring the unlinked DNA into a suitable recipient host cell.

- e) selecting and culturing the transformed host cell, and optionally
- f) isolating the desired DNA.

30

Genomic human DNA according to step b) of the process described above is isolated from suitable human tissue, preferably from human placenta or human foetal liver cells, according to methods known in the art. A genomic DNA library is constructed therefrom by limited digestion with suitable restriction endonucleases following established procedures. The genomic DNA library is replicated, e.g. on nitrocellulose membranes, and screened with a DNA probe for the DNA sequences of interest. The desired DNA may be amplified using PCR technology.

The transfer of the recombinant DNAs, e.g. the transfer of hybrid vectors, and the selection of transformed cells is described below.

Moreover, the invention relates to a suitable host cell transformed with the recombinant DNAs described above, namely a host cell which is transformed with a DNA encoding the light chain and/or a DNA encoding the heavy chain of the desired reshaped human antibody of the invention. It is preferred that the host cell contains a large number of copies of the vectors per cell.

The host cells of the present invention have to be capable of culture in vitro. Suitable host cells are of prokaryotic or eukaryotic origin and include bacterial cells, particularly E. coli, yeasts, e.g. Saccharomyces cerevisiae, or mammalian cells. To provide a suitable environment for the production of functional tetrameric antibodies, host cells of eukaryotic, particularly mammalian or yeast origin are preferred since the biosynthesis of functional tetrameric antibody molecules requires correct nascent polypeptide chain folding and assembly. Procaryotic hosts, especially E.coli, may be used for the production of antibody fragments of the invention, e.g. Fab- and Fv-fragments.

Examples of suitable hosts are microorganisms which are devoid of or poor in restriction or modification enzymes, such as bacteria, in particular strains of Escherichia coli, and yeasts, for example Saccharomyces cerevisiae.

Preferred host cells according to the invention are mammalian cells, e.g. COS-7 cells, Bowes melanoma cells, chinese hamster ovary (CHO) cells, embryonic lung cells L-132 and mammalian cells of lymphoid origin, such as lymphoma, myeloma, hybridoma, trioma or quadroma cells. Most preferred are mouse myeloma NSO cells.

These host cells are transfected with the light (L-) chain-gene construct alone, with the heavy (H-) chain-gene construct alone, or with both, either sequentially or simultaneously transferred with the help of two separatevectors or in a one-step procedure by using a double-construct (L-chain/ H-chain) vector as indicated hereinbefore. In the alternative, unlinked gene constructs may be transfected into the host cells either sequentially or simultaneously.

Preferred are host cells transfected with both gene constructs secreting reshaped human antibodies as described hereinbefore, particularly cell line EH31.8. Further examples of host cells of the invention are cells transfected with similar recombinant plasmids which contain alternative orientations of the H- and L-chain gene constructs, incorporating additional DNA elements to facilitate high levels of expression of the antibodies of the invention.

The host cells of the invention are genetically stable, produce and preferably secrete reshaped human antibodies of the invention of constant specificity and can be activated from deep-frozen cultures by thawing and recloning.

The transformed host cells are cultured by methods known in the art in a liquid medium containing assimilable sources of carbon, e.g. carbohydrates such as glucose or lactose, nitrogen, e.g. amino acids, peptides, proteins or their degradation products such as peptones, ammonium salts or the like, and inorganic salts, e.g. sulfates, phosphates and/or carbonates of sodium, potassium, magnesium and calcium. The medium furthermore contains, for example, growth-promoting substances, such as trace elements, for example iron, zinc, manganese and the like.

The medium is preferably chosen as to exert a selection pressure and prevent the growth of cells which have not been transformed or have lost the hybrid vector. Thus, for example, an antibiotic is added to the medium if the hybrid vector contains an antibiotic resistance gene as marker. If, for instance, a host cell is used which is auxotrophic in an essential amino acid whereas the hybrid vector contains a gene coding for an enzyme which complements the host defect, a minimal medium deficient of said amino acid is used to culture the transformed cells.

Culturing is effected by processes which are known in the art. The culture conditions, such as temperature, pH value of the medium and fermentation time, are chosen so that a maximum titer of the polypeptide or derivative of the invention is obtained. Thus, an <u>E. coli</u> or yeast strain is preferably cultured under aerobic conditions by submerged culture with shaking or stirring at a temperature of about 20°C to 40°C, preferably at about

30°C, and a pH value of 4 to 8, preferably of about pH 7, for about 4 to 30 hours, preferably until maximum yields of the polypeptide or derivative of the invention are reached.

When the cell density has reached a sufficient value, the culture is interrupted and the polypeptide or derivative can be isolated. If the hybrid vector contains a suitable secretion signal sequence, the polypeptide or derivative is secreted by the transformed cell directly into the culture medium. Otherwise, the cells have to be destroyed, for example by treatment with a detergent such as SDS, NP-40TM, TritonTM or deoxycholic acid, lysed with lysozyme or a similarly acting enzyme, or disrupted by an osmotic shock or ultra-sound. Break-up of the cells will also be required if the signal sequence directs the secretion of the desired protein into the cell periplasm. If yeast is used as a host microorganism, the cell wall may be removed by enzymatic digestion with a glucosidase. Alternatively or additionally, mechanical forces, such as shearing forces (e.g. French press, Dyno mill and the like) or shaking with glass beads or aluminium oxide, or alternating freezing, for example in liquid nitrogen, and thawing, for example at 30°C to 40°C, as well as ultra-sound can be used to break the cells.

The cell supernatant or the solution obtained after centrifugation of the mixture obtained after breaking the cells, which contains proteins, nucleic acids and other cell constituents, is enriched in proteins, including the polypeptides of the invention, in a manner which is known <u>per se</u>. Thus, for example, most of the non-protein constituents are removed by polyethyleneimine treatment and the proteins including the polypeptides and derivatives of the invention are isolated e.g. by the methods mentioned above.

The invention also relates to processes for the preparation of transformed host cells characterized in that suitable recipient host cells as described hereinbefore are transformed with one or two vectors according to the invention, and the transformed cells are selected.

Transformation of microorganisms is carried out as described in the literature, for example for <u>S. cerevisiae</u> (A. Hinnen et al., Proc. Natl. Acad. Sci. USA 75: 1929, 1978), and for <u>E. coli</u> (M. Mandel et al., J. Mol. Biol. 53: 159, 1970).

Accordingly, the transformation procedure of <u>E</u>. <u>coli</u> cells includes, for example, Ca²⁺ pretreatment of the cells so as to allow DNA uptake, and incubation with the hybrid vector. The subsequent selection of the transformed cells can be achieved, for example, by transferring the cells to a selective growth medium which allows separation of the transformed cells from the parent cells dependent on the nature of the marker sequence of the vector DNA. Preferably, a growth medium is used which does not allow growth of cells which do not contain the vector. The transformation of yeast comprises, for example, steps of enzymatic removal of the yeast cell wall by means of glucosidases, treatment of the obtained spheroplasts with the vector in the presence of polyethylene glycol and Ca²⁺ ions, and regeneration of the cell wall by embedding the spheroplasts into agar. Preferably, the regeneration agar is prepared in a way to allow regeneration and selection of the transformed cells as described above at the same time.

Transformation of cells of higher eukaryotic origin, such as mammalian cell lines is preferably achieved by transfection. Transfection is carried out by conventional techniques, such as calcium phosphate precipitation, microinjection into the cell nucleus, protoplast fusion, electroporation, i.e. introduction of DNA by a short electrical pulse which transiently increases the permeability of the cell membrane, or the like. Transfection may be carried out in the presence of helper compounds, e.g. diethylaminoethyldextran, dimethyl sulfoxide, glycerol, polyethylene glycol or the like, or as co-precipitates of vector DNA and calcium phosphate.

After the transfection procedure, transfected cells are identified and selected with the help of a selection procedure matching the selection marker of the DNA used for transfection. Selection markers include genes which confer resistance to heavy metals such as copper or to antibiotics, e.g. G-418 (geneticin, a neomycinderivative) or hygromycin, or genes which complement a genetic lesion of the host cell such as the absence of thymidine kinase, hypoxanthine phosphoribosyl transferase, dihydrofolate reductase, or the like. For example, if the DNA used for transfection comprises a marker for geneticin resistance, transformed cells are identified and separated from untransformed cells by culture in the presence of the antibiotic geneticin.

45

A reshaped human antibody according to the invention or a derivative thereof is useful for the qualitative and quantitative determination of IgE, especially in body fluids, e.g. in serum, in vitro and in vivo.

For instance, the reshaped human antibody or a derivative thereof can be used in any of the known immunoassays which rely on the binding interaction between the antigenic determinants of IgE and the paratopes of said antibody. Examples of such assays are radioimmunoassays (RIA), enzyme, immunofluoresence, chemiuminescence, immunoprecipitation, latex agglutination, or hemagglutination immunoassays.

The reshaped human antibody according to the invention can be used as such or in the form of radioactively labelled derivative in a radioimmunoassay (RIA). Any of the known modifications of a RIA can be used, for example soluble phase (homogeneous) RIA, solid phase (heterogeneous) RIA, single RIA or double (sandwich) RIA with direct or indirect (competitive) determination of IgE.

An example of such a radioimmunoassay is a sandwich RIA in which a suitable carrier, for xample the plastic surface of a microtitre plate or of a test tube, e.g. of polystyrene, polypropylene or polyvinylchloride,

glass or plastic beads, filter paper, dextran etc., cellulose acetate or nitrocellulose sheets, magnetic particles or the like, is coated with an antibody of the invention by simple adsorption or optionally after activation of the carrier. Then test solutions containing IgE and finally a reshaped antibody which also reacts with the antigen and which is radioactively labelled, e.g. with ¹²⁵I, is added. The amount of IgE in the test solutions is directly proportional to the amount of bound reshaped antibody and is determined by measuring the radioactivity bound to the carrier.

A reshaped human antibody according to the invention can be used as such or in the form of an enzymeconjugated derivative in an enzyme immunoassay. As described above for radioimmunoassays, any of the known modifications of an enzyme immunoassay can be used.

The tests are carried out in an analogous manner to the radioimmunoassays described above using an enzyme label instead of a radioactive label. The amount of immune complex formed which corresponds to the amount of IgE present in the test solutions is determined by adding an enzyme substrate solution. The enzyme substrate reaction results, for example, in a color change which can be observed by eye or with optical measuring devices.

A reshaped antibody according to the invention can be used as such or in the form of a derivative conjugated with chemiluminescent markers in a chemiluminescence assay. As described above for radioimmunoassays, any of the known modifications of a chemiluminescence assay can be used.

The tests are carried out in an analogous manner to the radioimmunoassays described above using a chemiluminescent label instead of a radioactive label. The amount of immune complex formed which corresponds to the amount of IgE present in the test solutions is determined by adding a compound triggering luminescence, e.g. H₂O₂ and NaOH, and measuring the emission of light with optical measuring devices.

The use according to the invention of a reshaped human antibody or a derivative thereof as described hereinbefore for the determination of IgE also includes other immunoassays known per se, for example immunofluorescence assays, latex agglutination with antibody-coated or antigen-coated latex particles, hemagglutination with antibody-coated or antigen-coated red blood corpuscles, evanescent light assays using an antibodycoated optical fibre and other direct-acting immunosensors which convert the binding event into an electrical or optical signal, or the like.

A reshaped human antibody according to the invention or a derivative thereof is also useful for the determination of IgE-producing cells, preferentially in a plaque forming cell (PFC) assay.

A plaque forming cell assay according to the invention is based on the principles of a solid phase immunoassay. Any of the known modifications of a solid phase immunoassay can be used, for example a radioimmunoassay, an enzyme, immunofluorescence or chemiluminescence immunoassay, or the like.

An example of such a plaque forming cell assay is a PFC assay based on an enzyme-linked immunosorbent assay (ELISA). For determination of the total amount of IgE-producing cells, a suitable carrier as described above for a sandwich RIA is coated with an antibody of the invention. A suspension of IgE-producing cells which are obtained from body fluids containing such cells by centrifugation, filtration, or the like, and a second polyclonal or monoclonal antibody specific for IgE, e.g. an antibody of the invention recognizing a different epitope of IgE than the first antibody, which is conjugated with an enzyme, e.g. alkaline phosphatase, are added. The amount of IgE-producing cells in the test suspensions is directly proportional to the amount of bound second antibody and is determined by adding an appropriate substrate solution, which results for example in the development of a colored reaction product, and counting the colored spots (plaques). For determination of the fraction of IgE-producing cells which produce IgE directed against a specific allergen, the carrier is first coated with the allergen or an adsorbable conjugate of the allergen before adding a cell suspension as described above. The fraction of IgE in the test suspension which is directed against the allergen binds to the surface-bound allergen and is determined by adding an antibody of the invention conjugated with an enzyme and an appropriate substrate solution resulting for example in the development of a coloured reaction product, and counting the colored spots (plaques).

Furthermore, the invention concerns test kits for the qualitative and quantitative determination of IgE and/or IgE producing cells comprising monoclonal antibodies and/or derivatives thereof of the invention and, optionally, other monoclonal or polyclonal antibodies and/or adjuncts.

Test kits according to the invention for a radioimmunoassay contain, for example, a suitable carrier, optionally freeze-dried or concentrated solutions of one or more monoclonal antibodies, solutions of a radioactively labelled monoclonal antibody or of radioactively labelled IgE, standard solutions of IgE, buffer solutions and, optionally, detergents for preventing non-specific adsorption and aggregate formation, pipettes, reaction vessels, calibration curves and the like. One or more of the monoclonal antibodies of the test kit are monoclonal antibodies of the invention. Test kits for the determination of IgE-producing cells which produce IgE directed against a specific allergen additionally contain solutions of the allergen or an adsorbable conjugate of the allergen.

Test kits according to the invention for an enzyme-immunoassay contain, for example, a suitable carrier, optionally freeze-dried or concentrated solutions of one or more monoclonal antibodies, optionally freeze-dried or concentrated solutions of an enzyme-labelled monoclonal antibody, of enzyme-labelled IgE, of a polyclonal anti-IgE serum and/or of enzyme-labelled monoclonal or polyclonal antibodies that recognize and bind the anti-IgE antibody, enzyme substrates in solid or dissolved form, standard solutions of IgE, buffer solutions, detergents, pipettes, reaction vessels, calibration curves, color scale tables and the like. One or more of the monoclonal antibodies of the test kit are monoclonal antibodies of the invention. Test kits for the determination of IgE-producing cells which produce IgE directed against a specific allergen additionally contain solutions of the allergen or an adsorbable conjugate of the allergen.

Moreover, the reshaped human antibodies according to the invention and their derivatives can be used for the qualitative and quantitative determination of surface IgE positive (sIgE*) B cells by any of the known conventional staining techniques, e.g. by flow cytometric analysis.

In addition, the monoclonal antibodies of the invention and/or their derivatives are useful for the treatment and/or prophylaxis of allergy.

The therapeutic effect is achieved by downregulating the IgE immune response due to the specific characteristics of the reshaped human antibodies and derivatives thereof according to the invention:

15

20

25

30

- They are capable of neutralizing formed IgE by binding free IgE and inhibiting the binding of IgE to cells bearing Fc, receptors I or II, in particular mast cells and basophils.
- They recognize and bind IgE expressed on the surface of surface IgE positive B cells (slgE+ B cells) and are therefore useful in depleting the population of such cells which form a "memory pool" resulting in IgE production after a second exposure to the allergen. The possibility of producing reshaped human antibodies of chosen immunoglobulin (sub)classes allows the activation of cellular mechanisms of the host immune system resulting in specific killing of the slgE+ B cells. This can also be achieved by conjugates of the monoclonal antibodies of the invention with cytotoxic drugs which will deliver such drugs to the target cells.
- Since the reshaped human antibodies of the invention and their derivatives do not recognize cytophilic lgE on cells bearing Fc_e receptors I or II, e.g. mast cells and basophils, they do not induce mediator release by these cells.
- The monoclonal antibodies and derivatives thereof according to the invention also have a long lasting therapeutic effect because they have a significant inhibitory effect on the formation of IgE in the immune response.

In consequence, the reshaped human antibodies of the invention and their derivatives provide a treatment that, rather than treating symptoms, actually affects the underlying cause of allergy, for example by removal of IgE antibodies and surface IgE positive B cells, thus eliminating the potential for an allergic response, and inhibition of IgE formation. It is especially advantageous that the treatment does not require ongoing repeated doses, and that the reshaped human antibodies and their derivatives of the invention can be used for prophylactic treatment by administration prior to detection of any of the symptoms of allergy.

As they are only weakly immunogenic or non-immunogenic when administered to humans the reshaped human antibodies and derivatives thereof according to the invention are especially useful for in vivo diagnostics, therapeutic applications and prophylaxis. Preferably, the reshaped human antibodies are tolerated by the human organism as self proteins when administered for therapeutic purposes.

The therapeutic daily dose for mammals is between approximately 0.1 mg and 10 mg per kg body weight depending on the status of the patient and the mode of application.

The Invention also relates to pharmaceutical preparations comprising a reshaped human antibody and/or derivatives thereof according to the Invention. The pharmaceutical preparations comprise, for example, the reshaped human antibodies and/or derivatives thereof in a therapeutically effective amount together or in admixture with inorganic or organic, solid or liquid pharmaceutical carriers.

Preferred are pharmaceutical preparations for parenteral application and inhalation. Preparations for intramuscular, subcutaneous or intravenous application or for inhalation are e.g. isotonic aqueous solutions or suspensions, optionally prepared shortly before use from lyophilized or concentrated preparations. The pharmaceutical preparations may be sterilized and contain adjuvants e.g. for conserving, stabilizing, wetting, emulsifying or solubilizing the ingredients, salts for the regulation of the osmotic pressure, buffer and/or compounds regulating the viscosity, e.g. sodium carboxycellulose, dextran, polyvinylpyrrolidone or gelatine. They are prepared by methods known in the art, e.g. by conventional mixing, dissolving or lyophilizing, and contain from approximately 0.01 % to approximately 50 % of active ingredients. The preparations for injections are processed, filled into ampoules, vials or disposable injection devices, and sealed under aseptic conditions according to methods known in the art.

The pharmaceutical preparations of the invention may be used for the prophylaxis and treatment of allergic

reactions in humans, in particular those typical of immediate type hypersensitivity as associated e.g. with allergic asthma, allergic rhinitis and atopic excema.

Brief description of the drawings:

Fig. 1: HCMV- mammalian expression vectors used to produce C21-L1 fused to the human κ light chain constant domain and C21-H1 fused to the human γ 1 heavy chain constant domain.

The invention particularly concerns the reshaped human antibodies, the recombinant DNAs, the transformed host cells, and the method for the preparation thereof as described in the Examples. The following examples illustrate the invention but do not limit it to any extent.

Abbreviations: V_L = light chain variable region; V_H = heavy chain variable region; CDR = complementarity determining region; FR = framework region; HCMV = Human Cytomegalovirus

Materials

15

Human IgGs with k light chains, purified from human plasma, are all purchased from Sigma, Buchs, Switzerland (IgG1: I-3889, IgG2: I-4139, IgG3: I-4389 and IgG4: 1-4639). Human IgM (Cat No. PHP003) and human IgD (Cat.No. PHP005) from human myeloma serum are obtained from Serotec. IgAs from human plasma (IgA1: 400105; IgA2: 400108) are from Calbiochem, Läufelfingen, Switzerland.

Example 1: Molecular modelling of mAb C21 V_L and V_H

A molecular model of the V_L (SEQ. ID NO. 1) and V_H (SEQ. ID NO. 3) regions of mouse monoclonal antibody C21, which recognizes human IgE, is built, for V_L , on the solved structure of the highly homologous mouse anti-lysozyme antibody HyHEL-10 (Padlan, E.A., Silverton, E.W., Sheriff, S., Cohen, G.H., Smith-Gill and Davies, D.R., 1989, Proc. Natl. Acad. Sci., USA, 86:5938; referred to as sequence 3 HFM in the Brookhaven Database, Bernstein et al., J. Mol. Biol 112, 535-542 (1977)) and, for V_H , on the structure of the mouse anti-lysozyme antibody HyHEL-5 (Sheriff, S., Silverton, E.W., Padlan, E.A., Cohen, G.H., Smith-Gill, S.J., Binzel, B.C. and Davies, D.R., 1987, Proc. Natl. Acad. Sci., USA, 84:8075; referred to as sequence 2 HFL in the Brookhaven Database, supra). The light and heavy chain variable regions of mAb C21 and HyHEL-10 or HyHEL-5 have 91% and 90% amino acid identity, respectively. The model is built on a Silicon Graphics IRIS 4D workstation running under the UNIX operating system and using the molecular modelling package QUANTA (Polygen Corp., USA). Identical residues in the framework are retained; non-identical residues are substituted using the maximal overlap procedure (Snow, M.E. and Amzel, L.M., 1986, Proteins 1:267) incorporated into QUANTA's protein modelling facility.

The complementarity determining regions CDR1 (L1), CDR2 (L2) and CDR3 (L3) of the V_L region and CDR1 (H1) and CDR2 (H2) of the V_H region from mouse C21 antibody correspond to canonical forms postulated previously (Chothia, C., Lesk, A.M., Tramontano, A., Levitt, M., Smith-Gill, S.J., Air, G., Sherrif, S., Padlan, E.A., Davies, D., Tulip, W.R., Colman, P.M., Spinelli, S., Alzari, P.M. and Poljak, R.J., 1989, Nature, 342:877). The main chain torsion angles of these loops are kept as in the original antibody structures (HyHEL-10 for L1-L3 and HyHEL-5 for H1-H2). There are no canonical structures for the CDR3 (H3) of the V_H regions, it is therefore modelled differently. Thirty candidate loops are extracted from 91 high resolution protein structures using a published algorithm (Jones, T.A. and Thirup, S., 1986, EMBO J., 5:819-822) as implemented in QUANTA, and the best version selected by eye. The loops are anchored on three framework residues on either side of the H3 CDR. Thus, H3 of the V_H region is modelled on Bence-Jones protein RHE (Furey, W., Wang, B.C., Yoo, C.S. and San, M., 1983, J. Mol. Biol., 167:661-692) in the region of residues 87-106, which corresponds roughly to CDR3 (L3).

The model is subjected to steepest descents and conjugate gradients energy minimization using the CHARM potential (Brooks, B.R., Bruccoleri, R.E., Olafson, B.D., States, D.J., Swaminathan, S. and Karplus, M., 1983, J. Comp. Chem., 4:187) as implemented in QUANTA in order to relieve unfavourable atomic contacts and to optimize Van der Waals and electrostatic interactions.

Example 2: Design of reshaped human C21 V_L and V_H regions

The design of reshaped human C21 V_L and V_H regions is based primarily on the consensus sequences of human V_L and V_H regions (versions C21-L1, C21-L2, C21-L3, C21-H1 and C21-H3) as found in the KABAT database (Kabat, E.A., Wu, T.T., Reid-Miller, M., Perry, H.M. and Gottesman, K.S., 1987, Sequences of Proteins of Immunological Interest, 4th Edition, U.S. Department of Health and Human Services, U.S. Government

Printing Office). In addition, two more reshaped human C21 V_H regions (C21-Hay1 and C21-Hay3) are based on the framework regions (FRs) of an individual human antibody.

For the design of consensus-based reshaped human C21 variable regions, the amino acid sequences of the V_L and V_H regions from mouse C21 antibody are compared with the consensus sequences for V_L and V_H regions of human antibodies from the KABAT database. This analysis reveals that the mouse C21 V_L region and the mouse C21 V_H region are most similar to the human κV_L subgroup III consensus sequence (77% amino acid sequence identity), respectively. These human consensus sequences are used to design the reshaped human C21 light and heavy chain variable regions C21-L0 and C21-H0 containing murine C21 CDRs and the human FRs of the respective consensus sequence. The molecular models of the mouse C21 variable regions (Example 1) are used to identify framework residues that are potentially important to achieve good antigen binding and which might be critical for V_L/V_H packing. As a result of this graphical analysis, some of the noted human consensus amino acids within the FRs are exchanged for their corresponding mouse C21 residues. These changes are only considered within the human framework region if they do not fall into one of the following categories:

[1] The human consensus sequence (human subgroup 1; HSG1) reveals no dominant amino acid preference at this position, but the amino acid as found in the original mouse C21 sequence is present in at least one individual sequence of the respective human immunoglobulin variable region subgroup. For example, HSG 1 is described as having no consensus sequence at amino acid residue 19 (Kabat et al., supra) although several human antibodies have a Lys residue at this position. Since Lys also appears in the C21 sequence it is retained in the reshaped monoclonal antibody.

[2] The amino acid at the framework position is part of a postulated canonical structure, important in determining the structure of the CDRs or hypervariable loops, and is thus expected to be indispensible for maintaining the shape and integrity of the antigen binding site (Chothia, C. and Lesk, A.M., 1987, J. Mol. Biol., 196:901; Chothia, C. et al., 1989, supra).

According to these rules, four and six amino acids within the reshaped human light and heavy chain variable regions C21-L0 and C21-H0 are exchanged when compared with the human consensus sequences, resulting in versions C21-L1 and C21-H1. The positions of the exchanged amino acids are 1, 3, 49 and 60 in C21-L1 and its below-identified modified version C21-L1 (SEQ ID NO. 5). In C21-H1 (SEQ. ID NO. 11) the positions of the exchanged amino acids are 38, 40, 67, 68, 70 and 87. An exception to the above-mentioned rules is position 76 of the reshaped human C21 V_H region, where we choose the most frequent human amino acid for this position (Thr) as found in the human V_H consensus sequence.

Further new versions of V_L and V_H contain the following alterations (compared with C21-L1 and C21-H1, respectively):

C21-L2 (SEQ ID NO. 7): aspartic acid (instead of serine) at position 60;

20

25

35

C21-L3 (SEQ ID NO. 9): glutamic acid (instead of aspartic acid) at position 1; valine (instead of leucine) at position 3:

C21-H3 (SEQ ID NO. 13): arginine (instead of lysine) at position 38; alanine (instead of arginine) at position 40; arginine (instead of lysine) at position 67; arginine (instead of threonine) at position 87.

A database search using C21-L1' and C21-H1 reveals that reshaped human C21 V_L version C21-L1' is most similar (91% sequence identity) to human κ light chain variable region HUMIG KAF (EMBL database, Heidelberg, Germany; Newkirk, M.M., Gram, H., Heinrich, G.F., Oestberg, L., Capra, J.D. and Isserman, R.L., 1988, J. Clin. Invest., 81:1511-1518) and that reshaped human C21 V_H version C21-H1 is most similar (78% sequence identity) to human heavy chain variable region HUMIG HAY (EMBL database, supra; Dersimonian, H., Schwartz, R.S., Barrett, K.J. and Stollar, B.D., 1987, J. Immunol., 139:2496-2501). These sequences are referred to below as KAF and HAY, respectively. The FRs of KAF and the human κ V_L subgroup III consensus sequence differ only at positions 49 and 85. At position 49 the corresponding mouse C21 amino acid (lysine) is retained, due to its putative antigen binding, whereas at position 85 the human V_L III consensus amino acid valine is changed to methionine, as found in KAF. Thus, the modified version of C21-L1', designated C21-L1 (SEQ. ID. NO. 5), is based on the individual human light chain variable region KAF (Newkirk, M.M. et al., 1988, supra). It differs from the first version C21-L1' in that there is a methionine instead of a valine at position 85. The reshaped versions C21-L2 (SEQ ID NO. 7) and C21-L3 (SEQ ID NO. 9) also have a methionine at position 85.

In contrast, the FRs of human heavy chain variable region HAY and the human V_H subgroup I consensus sequence differ at several positions. In order to construct reshaped human C21 V_H regions that show a high degree of similarity to this individual human antibody, two more versions of reshaped human C21 V_H regions are designed based on the FRs from the heavy chain variable region of human heavy chain variable region HAY. The construction of the HAY-based reshaped human C21 V_H versions necessitate five changes in FR2 and FR3 of reshaped human C21 versions C21-H1 and C21-H3 at positions 43, 44, 48, 76 and 77, respectively.

These HAY-based versions are called C21-Hay1 (SEQ ID NO. 15) and C21-Hay3 (SEQ. ID NO. 17), respectively. Two additional differences between the FRs of HAY and the human V_H subgroup I consensus sequence at positions 30 and 72 are retained as in the mouse C21 V_H region, and are not regarded as changes, because they are canonical residues that define the structure of the CDRs (Chothia, C. et al., 1989, supra).

Example 3: Design and Construction of Humanized Antibody Genes

· For the design of humanized antibody gene cassettes, additional sequences necessary for efficient expression and cloning are added at the 5'- and 3'-ends of the resulting coding regions. Eukaryotic leader sequences for efficient expression of reshaped human C21 antibodies are added in frame to the designed humanized variable regions. The leader sequence for the reshaped human C21 V₁ region is derived from the leader sequence found in the k light chain of human antibody KAF (Newkirk, M.M. et al., 1988, supra) from which the variable region is used for reshaping of the C21 V₁ region. The leader sequence for the reshaped human C21 VH region is derived from the leader sequence found in the heavy chain of human antibody HG3 CL (Rechavi, G., Ram, D., Glazer, L., Zakut, R. and Givol, D., 1983, Proc. Natl. Acad. Sci., USA, 80:855-859), a member of human VH subgroup I (Kabat, E.A., Wu, T.T., Reid-Miller, M., Perry, H.M. and Gottesman, K.S., 1987, Sequences of Proteins of Immunological Interest, 4th Edition, U.S. Department of Health and Human Services, U.S. Government Printing Office). The resulting protein sequences for reshaped human C21 VL/VH regions are then back-translated into DNA sequences using the Codon Usage Table for mouse sequences, as found in the Sequence Analysis Software Package of the Genetics Computer Group, University of Wisconsin, USA. To the designed DNA fragments are also added eukaryotic translation signals at the 5'-end (Kozak, M., 1987, J. Mol. Biol., 196: 947-950), donor splice sites at the 3'-end (Breathnach, R., Benoist, C., O'Hare, K., Gannon, F. and Chambon, P., 1978, Proc. Natl. Acad. Sci,. USA, 75:4853-4857) and Hind III (5'-ends) and Barn HI and Xba I (3'-ends) DNA restriction sites for convenient subcloning into the designated mammalian expression vectors.

The designed humanized antibody gene cassettes, encoding the reshaped human C21 V_L and V_H regions C21-L1 and C21-H1 are then constructed by gene synthesis using synthetic DNA polynucleotides. The entire DNA fragments are subdivided into six regions overlapping with each other by 20 nucleotides. For each reshaped human C21 variable region gene cassette, six 5'-phosphorylated and PAGE-purified polynucleotides designated C21-LA (SEQ ID NO: 19), C21-LB (SEQ ID NO: 20), C21-LC (SEQ ID NO: 21), C21-LD (SEQ ID NO: 22), C21-LE (SEQ ID NO: 23), C21-LF (SEQ. ID NO: 24), C21-HA (SEQ ID NO: 25), C21-HB (SEQ ID NO: 26), C21-HC (SEQ ID NO: 27), C21-HD (SEQ ID NO: 28), C21-HE (SEQ ID NO: 29) to HF (SEQ. ID NOs. 30) are purchased from Genosys Biotechnologies, Houston, Texas, USA. They are then assembled in a polymerase chain reaction (PCR)-based gene synthesis. 5 pmol of each polynucleotide (i.e. LA to LF) are first annealed and extended in a 100 µl reaction containing 10 mM Tris-HCl pH 8.3, 1.5 mM MgCl₂, 50 mM KCl, 10 mM β-mercaptoethanol, 0.05% (w/v) Tween-20, 0.05% NP-40 (Merck, Zürich), 200 μM dNTPs (N = G, A, T or C) and 5 U VentTM DNA polymerase (New England Biolabs). Temperature steps are 95°C/1 min, 50°C/2 min and 72°C/4 min using a Techne PHC-2 temperature cycler. After this first cycle, 50 pmol of oligonucleotide primers C21-5' (SEQ ID NO. 31) and C21-L3' (SEQ ID NO. 33) or C21-H3' (SEQ ID NO. 32) hybridizing at the 5'- and 3'-ends of the desired, full-length DNA fragment are added and the full-length DNA fragment amplified in a PCR reaction of about 20 cycles using the following cycling parameters: 95°C/1 min, 60°C/2 min and 72°C/2 min. The PCR mixture is then extracted once with one volume of chloroform and the DNA precipitated by adding 1/10 vol of 8 M LiCl and 3 vol of ethanol. The precipitated DNA is redissolved in H₂O and digested with Hind III and Barn HI restriction endonucleases under conditions suggested by the supplier (Boehringer, Mannheim, Germany). DNA fragments of the expected sizes (C21-L1 = 416 bp and C21H1 = 459 bp) are electrophoretically purified in 1% agarose/TBE and excised from the gel. Gel pieces are cut into smaller fragments, frozen in liquid nitrogen for 5 min, and then eluted through glasswool by centrifugation in a microcentrifuge (30 min, 136000 x g). After phenol-chloroform extraction and LiCl/ethanol precipitation at room temperature, the purified Hind III-Bam HI restriction fragments are subcloned into pBluescript KS II M13+ (Stratagene) and transfected into competent E. coli cells (HB101 strain from GIBCO-BRL). Multiple plasmid clones of either KS+C21/L1 or KS+C21/H1 containing DNA inserts of correct size are sequenced using Sequenase (USB). Point mutations and/or deletions within the DNA sequence are corrected by exchanging DNA restriction enzyme fragments between different clones and/or oligonucleotide directed PCR mutagenesis according to a published procedure (Kammann, M., Laufs, J., Schell J. and Gronenborn B., 1989, Nucl. Acids Res., 17:5404). Hind III-Bam HI fragments exhibiting correct DNA sequences are then subcloned into the light or heavy chain expression vectors to create the plasmids HCMV-k-C21/L1 and HCMV-y1-C21/H1 (depicted in Fig. 1) wherein the Hind III-BamHI fragment coding for the reshaped human light or heavy chain variable region is joined to a DNA coding for human k and y1 constant regions, respectively. Both plasmids comprise the origin of Simian Virus 40 (SV40), th HCMV enhancer domain, the HCMV promoter, and the ampicillin selectable gene (Kettleborough et al., 1991, supra).

Reshaped human C21 V_L region versions C21-L2 and C21-L3 are generated by oligonucleotide-directed mutagenesis, making use of recombination events which occur during PCR reactions (Mullis, K., Faloona, F., Scharf, S., Saikl, R., Horn, G. and Ehrlich, H., 1986, Cold Spring Harbour Symp., 51:263 and Yolov, A.A. and Shabarova, Z.A., 1990, Nucl. Acids Res., 18:3983). Oligonucleotide primers C21-5'(SEQ ID NO. 31), L/D60SL (SEQ ID NO. 35), L/D60S-SL (SEQ ID NO. 36) (for C21-L2), RSP (SEQ ID NO. 34), L/E1D-V3L (SEQ ID NO. 37), L/E1D-V3L-SL (SEQ ID NO. 38), (for C21L3) and C21-L3' (SEQ ID NO. 33) (for C21-L2 and -L3) are synthesized in order to generate by PCR amplification two DNA fragments, for each of the two light chain V-region versions. Except for the terminal oligonucleotide primers (C21-5', RSP and C21-L3') oligonucleotides incorporate sequences needed for the desired codon changes. Except for primer pair RSP and L/E1D-V3L, about 50 pmol each of the appropriate primer pair is combined with ca. 10 ng of a Xho I-Not I fragment from KS+C21/L1, 3 units of VentTM DNA polymerase and 25 PCR amplification cycles are used (60°C/25 s, 72°C/40 s and 93°C/25 s). For primer pair RSP and L/E1D-V3L, ca. 150 ng of KS+C21/L1 plasmid DNA is used as a template and the desired DNA fragment PCR amplified using 35 cycles (40°C/30 s, 72°C/1 min and 93°C/30 s). The products of these reactions, purified by agarose gel electrophoresis, are first combined, and ca. 5-30 ng of each DNA fragment extended with 5 units of VentTM DNA polymerase (95°C/1 min, 50°C/2min and 72°C/4 min). Terminal oligonucleotide primers C21-5' and C21-L3' are then added and the combined full-length DNA fragment PCR amplified using 25 cycles (95°C/1 min, 50°C/2 min and 72°C/2 min). The PCR amplified DNAs for C21-L2 and C21-L3 are then subcloned into pBluescript KS II M13+ (Stratagene) after digestion using DNA restriction endonucleases Hind III and BamHI, and sequenced. Correct sequences are then transferred into the HCMV-y1-expression vector as described above.

Reshaped human C21 V_H region versions C21-H3, C21-Hayl and C21-Hay3 are generated in a similar manner as described for C21-L1 and C21-L2. To obtain these C21 V_H region versions, oligonucleotide primers C21-5', H/R38K-A40R-L (SEQ ID NO. 39), H/R38K-A40R-SL ((SEQ ID NO. 40), H/R67K-L (SEQ ID NO. 41), H/R67K-SL (SEQ ID NO. 42), H/R87T-L (SEQ ID NO. 43), H/R87T-S (SEQ ID NO. 44), HayFR2 (SEQ ID NO. 45), HayFR2-S (SEQ ID NO. 47), HayFR3 (SEQ ID NO. 48), HayFR3S (SEQ ID NO. 49) and C21-H3' are used to PCR amplify, using C21-H1 (for C21-H3 and C21-Hay1) or C21-H3 (for C21-Hay3) as DNA templates, double-stranded DNA fragments containing the desired codon changes for the different reshaped human C21 V_H versions. Corresponding, agarose gel-purified fragments are then assembled by PCR recombination to yield the full-length DNA-fragments, as described above. After digestion with DNA restriction endonucleases Hind III and Barn HI, followed by cloning into pBluescript KS II M13+ (Stratagene) as described above, plasmid clones are checked for the correct sequence before being cloned into the HCMV-y1-expression vector.

Example 4: Transient Expression of recombinant plasmids in COS cells

COS cells are electroporated using 10 μ g of the HCMV expression vectors bearing the genes coding for the reshaped human C21 heavy and light chains. 10 μ g of H- and L-chain expression plasmids are added to 0.8 ml of a 1 x 10⁷ cells/ml suspension of COS cells in PBS/o (PBS lacking Ca2+ and Mg2+ supplied by GIBCO-BRL, Basel, Switzerland; cat.no. 041-04190M). ABio-Rad Gene Pulser is then used to deliver to the suspended cells a pulse of 1900 V at a capacitance of 25 μ F. The cells are allowed to recover for 10 min before plating into 10 ml DMEM containing 5% v/v gammaglobulin-free and heat-inactivated fetal calf serum (GIBCO-BRL, Basel, Switzerland; cat. no. 063-06510H). After 72 h incubation, the medium is collected, centrifuged to remove cells and cellular debris. The COS cell supernatant is then filtered through a 0.45 μ m membrane and analysed for the presence of assembled antibody of human γ 1/ κ isotype by ELISA. The humanized antibodies are further purified by protein A affinity chromatography.

Example 5: ELISA assay for human IgG/k production

50

96-well microtiter plates (Nunc MaxiSorb, cat. No. 439454) are coated overnight with 50 μ l of a 1:1000 dilution of goat anti-human IgG (Fc specific; Dianova, #109-005-098) in PBS/o, pH 7.2. After this and all subsequent steps, plates are washed 3x with 200 μ l PBST (PBS/o pH 7.2 containing 0.05% Tween-20). Free binding sites are blocked for 1 h at 37°C with 100 μ l RIA-buffer (1% bovine serum albumin in PBST). 50 μ l of samples, and dilutions thereof in RIA-buffer, are added and the mixtures incubated for 1 h at 37°. Highly purified recombinant human antibody F5-444 (human γ 1/ κ isotype, European Patent Application No. 498767) serves as a standard. 50 μ l of a 1:1000 dilution of affinity-purified goat anti-human κ -light chain antiserum conjugated with horseradish peroxidase (Sigma, Buchs, Switzerland; cat. no. A-7164) in RIA-buffer is then applied and incubated for 1 h at 37°C. 100 μ l of ABTS (2,2'-Azino-bis(3-ethylbenzthiazoline-6-sulfonic acid)) substrate sol-

ution (BioRad, Glattbrugg, Switzerland; #172-1064) are used for development. After an appropriate incubation time the enzymic reaction is stopped using an equal volume (100 µl) of 2% (w/v) oxalic acid. The absorption at 415 nm is used for quantitation of bound and fully assembled human antibody.

Example 6: Protein A Purification of Humanized Antibodies from COS Cell Supernatants

Transiently-expressed humanized antibodies are purified by affinity chromatography on a lml Prosep A column (Bioprocessing Ltd, Durham, England) packed into a HR 5/5 FPLC column (Pharmacia, Uppsala, Sweden). The column is run at constant flow rate of 2mi/min on an FPLC system (Pharmacia, Uppsala, Sweden) and protein eluting from the column is detected in a flow cell by u.v.-absorbance at 280nm. The column is prepared by washing with 10 column volumes of PBS/o pH 8.0 (20 mM NaPhosphate, 150 mM NaCl), pre-elution with 10 column volumes of 100 mM sodium citrate buffer pH 3.0 and re-equilibration with 10 column volumes of PBS/o pH 8.0. COS cell supernatants (20-50ml), clarified by filtration through a 0.45 μm membrane, are loaded directly on to the column with a peristaltic pump. The column is then washed with PBS/o pH 8.0 until the u.v.-absorbance returns to baseline. Bovine IgG is then eluted by washing with 100 mM sodium citrate buffer pH 5.0 until the baseline returns to zero. Finally humanized antibodies are eluted using 100 mM sodium citrate buffer pH 3.0 and the eluates immediately adjusted to pH 7.0 by addition of 1M Trizma-Base (Sigma, Buchs, Switzerland). The purity of the humanized antibodies is analysed by SDS-Polyacrylamide gel electrophoresis using Coomassie blue staining (Laemmli, U.K., 1970, Nature, 227:680-685). For biosensor analysis the neutralized Protein A eluates are concentrated in a Centricon-10 microconcentrator (Amicon) and the buffer changed to PBS/o pH 7.2. Depending on the purity of the antibody preparation, the protein concentration is quantified either by u.v.-absorption at 280 nm or by human y/k ELISA using a known, purified recombinant chimeric antibody F5-444 of matched isotype as a standard.

Example 7: Analysis of the Avidity and Specificity of mouse and reshaped human C21 antibodies by Biospecific Interaction Analysis (BIA)

The avidity and specificity of the different combinations of reshaped human C21 variable light and heavy chains are analyzed using real-time biospecific interaction analysis (Jönsson, U., Fagerstam, L., Ivarsson, B., Johnsson, B., Karlsson, R., Lundh, K., Löfas, S., Persson, B., Roos, H., Rönnberg, I., Sjölander, S., Stenberg, E., Stahlberg, R., Urbaniczky, C., Östlin, H. and Malmqvist, M., 1991, BioTechniques, 11:620-627). All experiments are performed on the BIAcore™ system (Pharmacia Biosensor AB, Uppsala, Sweden) using CM5 sensor chips. As capture antibodies, ca. 11,000 RU (11 ng/mm2) of polyclonal rabbit anti-mouse IgGI (Pharmacia Biosensor AB, Uppsala, Sweden, cat.no. BR-1000-55) or rabbit anti-human IgG (obtained from Pharmacia Biosensor AB) are immobilized on to the sensor chip surface using their amino groups and EDC/NHS chemistry essentially as described previously (Jönsson, U. et al., 1991, supra). Four experimental cycles are performed for each antibody to determine the association rate of binding to human IgE. Each cycle consists of binding of a constant amount of test antibody to the respective catching antibody, followed by the interaction of this test antibody with a fixed concentration of antigen (human IgE; monoclonal antibody SE44; 3.125, 6.25, 12.5 and 25 nM) followed by a final regeneration of the surface using 40 mM HCl. Experimental details are as follows:

[1] The flowrate is 5 µl per min;

45

- [2] HBS (10 mM Hepes, 3.4 mM EDTA, 150 mM NaCl, 0.05% BlAsurfactant, pH 7.4) is used as running buffer;
- [3] Test antibodies (in PBS/o pH 7.2) are diluted in HBS to a final concentration of 5-10 μg/ml, and bound to the capture antibody to obtain 1300-2200 RU (1.3-2.2 ng/mm2) of bound test antibody;
 - [4] Human monoclonal IgE (SE44) is passed over the bound test antibody for 9 min;
 - [5] 4 μ 40 mM HCI is used to remove antibody-antigen complexes and prepare the surface for the next cycle;
 - [6] The assay temperature is 25°C.
- The association constants of the antibody-antigen interactions are then calculated using computer programs implemented in the Biocore™ system.

For the determination of dissociation rate constants a similar protocol is used, except that a dissociation phase is included. Assay conditions are as described above. Test antibodies are first bound to the sensor chip surface via immobilised catching antibodies. IgE (SE44) at the highest concentration (25 nM) is allowed to bind to the antibody. Following binding HBS buffer is passed over the sensor chip surface at a constant flow rate of 5 µl/min and the decrease in resonance signal monitored over a period of 15 to 25 min. The sensor chip is finally regenerated by washing with 4 µl of 40 mM HCl solution. Since the dissociation of antibody: IgE complexes is a first-order reaction the linear parts of the sensorgramms are taken to calculate the dissociation

rate constants using computer progammes implemented in the BIACORE™ system.

The kinetic constants k_{ess} (velocity constant of the antibody-antigen association) and k_{diss} (velocity constant of the dissociation of the antibody-antigen complex) and avidity (represented by the equilibrium constant K_{eff}) of reshaped human C21 antibodies are summarized in Table 1.

Table 1: Kinetic constants and avidity of reshaped human C21 antibodies. For k_{ass} the number of independent experiments is given in brackets (n); each k_{diss} is determined in two independent experiments.

antibody	k _{ess} x 10 ⁵ M ⁻¹ s ⁻¹	k _{diss} x 10 ⁻⁵ s ⁻¹	K _{en} x 1010M-1
TES-C21	2.4±0.3 (6)	2.6±0.0	0.92±0.12
C21-H1/L1	2.6±0.1 (3)	7.0±1.1	0.87±0.32
C21-H1/L2	2.8±0.1 (3)	5.7±0.2	0.49±0.02
C21-H1/L3	2.9±0.4 (3)	6.2±1.0	0.47±0.10
C21-H3/L1	2.5±0.3 (3)	1.9±0.6	1.32±0.44
C21-H3/L2	2.5±0.3 (4)	4.4±0.7	0.57±0.11
C21-H3/L3	2.6±0.5 (3)	3.5±0.2	0.74±0.15
C21-Hay3/L1	2.5±0.3 (3)	4.1±0.6	0.61±0.12
C21-Hay3/L2	2.5±0.3 (3)	3.1±0.6	0.81±0.18
C21-Hay3/L3	2.6±0.3 (3)	15.9±1.8	0.16±0.03

All of these reshaped human C-21 antibodies have quite similar rates of association. The reductions in the avidity of binding are caused mainly by higher rates of dissociation. Different versions of reshaped human C21 light chains are analyzed for the importance of positions 1 and 3 at the amino terminus. Good binding to antigen is obtained when the amino acids at these positions are the same as those present in the C21 light chain. Using the entire human KAF FR1 decreases the binding regardless of the heavy chain partner.

The reshaped human C21 antibodies are also tested for binding to all isotypes of human immunoglobulins by biospecific interaction analysis. First, the test antibodies are separately bound to immobilized capture antibodies on sensor chips as described above. Cross-reactivity of these immobilized rabbit anti-human IgG and rabbit anti-mouse IgG1 antibodies is then blocked using chimeric anti-CEA antibody 10 μg/ml ReK.41 for 5 min (human γ1/κ, European Patent Application No. 323806), before 5 μg/ml of human immunoglobulins of all isotypes IgM, IgD, IgA1, IgA2, IgG4, IgG3, IgG2, IgG1 and IgE (SE44) are passed successively over this pretreated surface for 5 min each. Finally, the surface is regenerated with 40 mM HCI. Flowrates, temperature and other conditions are the same as described above. Sensorgramms are recorded with the BIAcoreTM system. The reshaped human C21 antibodies are specific for human IgE isotype.

Example 8: Making Permanent Cell Lines

15

25

30

45

Plasmid DNAs of reshaped TESC-21 human CMV H- and L- chain expression vectors for transfection are purified by centrifugation to equilibrium in caesium chloride gradients two times. The recombinant immunoglobulin genes are introduced into mouse myeloma NSO cells by electroporation with the use of a Gene Pulser apparatus (BioRad, Richmond, CA). NSO cells (2 to 3 x 10²) are washed with phosphate-buffered saline (PBS), and resuspended in 0.8 ml of PBS containing 10 μg each of BspCI-linearized H- and L- chain plasmid DNAs. Electroporation is performed at an electric field of 210 V and a capacitance of 960 μFD. Cells recovered are diluted in Protein-Free Hybridoma Medium (PFHM, Gibco) containing 2% FBS (fetal bovine serum), and plated out in 96-well microtiter plates at 104 cells per well. After 48 h incubation the transfectants are selected in PFHM containing 0.7 mg/ml G418 (Gibco) and 2% FCS (foetal calf serum). After 2 weeks, wells containing drug resistant colonies are screened for the production of human IgG by ELISA. An ELISA is developed for the quantitation of the recombinant human IgG/kappa antibody expressed in the culture supernatant. Immulon 2 (Dynatech Labs) 96-well plates are coated overnight with 100 μl of 0.5 μg/ml goat anti-human kappa antibody (Southern Biotech) in PBS at room temperature. Wells are then blocked with Blotto (5% dry milk powder in

PBS) for 1 h and washed with PBS containing 0.05% Tween-20 (PBST). Culture supernatant (50 µl) is added to the coated wells and incubated for 1 h. After washing with PBST, 100 µl of horseradish peroxidase-conjugated goat anti-human IgG(Fc) antibody (Jackson ImmunoResearch Lab), diluted at 1/50,000 in Blotto, is added to each well, and the plate is incubated for 1 hour. Peroxidase substrate solution containing 0.1% 3', 3", 5', 5"-tetramethyl benzidine (Sigma) and 0.003% Hydrogen peroxide (Sigma) is added at 100 µl per well and incubated at room temperature for 0.5 hour. The reaction is stopped by the addition of 50 µl of 2 M sulphuric acid and the O.D. of the reaction mixture in each well is read at 450 nm with a Dynatech MR5000 plate reader. Reshaped TESC-21 human CMV expression plasmids, H1, H3, L1, and L3, are used to transfect NSO cells in four combinations: H3L1, H3L3, H1L1, and H1L3. A total of 142, 122, 68, and 64 wells of H3L1, H3L3, H1L1, and H1L3, respectively, give O.D. readings of greater than 0.05 with background reading below 0.01. For each combination, one cell line is chosen based on IgG secretion levels: EH31.8, secreting the H3L1 antibody; EH33.16, secreting the H3L3 antibody; EH11.13, secreting the H1L1 antibody; and EH13.5, secreting the H1L3 antibody.

15 Deposition data:

25

30

35

45

55

The following cell lines have been deposited with the American Type Culture Collection (ATCC), 12301 Parklawn Drive, Rockville, MD 20852, U.S.A., on September 23, 1992 accession nos. given in brackets). Cell line EH31.8 producing reshaped human antibody H3L1 (HB 11130)

Cell line EH11.13 producing reshaped human antibody H1L1 (HB 11132)
Cell line TES-C21 producing murine monoclonal antibody TES-C21 (HB 11133)
Cell line EH33.16 producing reshaped antibody H3L3 (HB 11131)
Cell line EH13.5 producing reshaped antibody H1L3 (HB 11134).

SEQUENCE LISTING

(1)	GENERAL	INFORMATION

(i)	APPI	T.TC	MT-

15

20

30

35

- (A) NAME: CIBA-GEIGY AG
- (B) STREET: Klybeckstr. 141
- (C) CITY: Basle
 - (E) COUNTRY: Switzerland
 - (F) POSTAL CODE (ZIP): 4002
 - (G) TELEPHONE: +41 61 69 11 11
 - (H) TELEFAX: + 41 61 696 79 76
 - (I) TELEX: 962 991
 - (A) NAME: TANOX BIOSYSTEMS, INC.
 - (B) STREET: 10301 Stella Link
 - (C) CITY: Houston
 - (D) STATE: Texas
 - (E) COUNTRY: United States of America
 - (F) POSTAL CODE (ZIP): 77025
- (ii) TITLE OF INVENTION: Reshaped monoclonal antibodies against an immunoglobulin isotype
- (iii) NUMBER OF SEQUENCES: 49
- (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)

	(2) INFORMATION FOR SEQ ID NO:1:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 370 base pairs (B) TYPE: nucleic acid	
	(C) STRANDEDNESS: double (D) TOPOLOGY: linear	
15	(ii) MOLECULE TYPE: DNA (genomic)	
20	<pre>(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1369 (D) OTHER INFORMATION: /product= *heavy chain variable</pre>	
25		
٠.	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
30	CAG GTT CAG TTG CAG CAG TCT GGA GCG GAG CTG ATG AAG CCT GGG GCC Gln Val Gln Leu Gln Gln Ser Gly Ala Glu Leu Met Lys Pro Gly Ala 1 5 10 15	48
		96
35	TCA GTG AAG ATC TCC TGC AAG ACT ACT GGC TAC ACA TTC AGT ATG TAC Ser Val Lys Ile Ser Cys Lys Thr Thr Gly Tyr Thr Phe Ser Met Tyr 20 25 30	.50
40	TGG TTA GAG TGG GTA AAG CAG AGG CCT GGA CAT GGC CTT GAG TGG GTT	144
	Trp Leu Glu Trp Val Lys Gln Arg Pro Gly His Gly Leu Glu Trp Val 35 40 45	
45	GGA GAG ATT TCA CCT GGA ACT TTT ACT ACT AAC TAC AAT GAG AAA TTC Gly Glu Ile Ser Pro Gly Thr Phe Thr Thr Asn Tyr Asn Glu Lys Phe 50 55 60	192

5	AAG	GCC	AAG	GCC	ACA	TTC	ACT	GCG	GAT	ACA	TCC	TCC	AAC	ACA	GCC	TAC		240
	Lys	λla	Lys	Ala	Thr	Phe	Thr	Ala	Asp	Thr	Ser	Ser	Asn	Thr	Ala	Tyr		
	65					70					75					.80		•
10	CTC	CAA	CTC.	ACC	GGC	CTY:	ACA	TCT	GAG	GAC	ՎԻ ԴԻ	ccc	CTC	тас	ጥጥ	TĠT		288
								Ser										
		•			85					90				-	95	-		
15	•						,											
13	GCA	AGA	TTC	TCC	CAT	TTT	TCC	GGT	AGT	AAC	TAC	GAC	TAC	TIT	GAC	TAC		336
	Ala	Arg	Phe		His	Phe	Ser	Gly		Asn	Tyr	Asp	Tyr		Asp	Tyr		
				100					105					110				
20	TGG	GGC	CAG	GGC	ACC	TCT	CTC	ACA	GTC	TCC	TCC	G						370
								Thr									. ;	
		•	115					120										
25																		
•	(2)	-	001/3	mr () 1	500	600	75.											
	(2)	INF	OKMA	TION	FUR	SEQ	In 1	NO:2	:									
 30			(i)	SEQU	ENCE	СНА	RACT	ERIS	TICS	:								
			(.	A) L	ENGT	н: 1	23 a	mino	aci	đs								
			(B) T	YPE:	ami	no a	cid										
•			(-	D) T	OPOL	OGY:	lin	ear										
35																		
		(ii) MO	LECU	LE T	YPE:	pro	tein										
		(xi) SE	OUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:2:							
40				_					_									
	Gln	Val	Gln	Leu	Gln	Gln	Ser	Gly	Ala	Glu	Leu	Met	Lys	Pro	Gly	Ala		
	1				5					10					15	i		
	_			-1-		_	•	-				. m\	nh a	0				
45	Ser	vai	Lys	20		Cys	Lys	rnr	25		туг	·THE	Pne	Ser 30		Tyr		
				20	•				2,3					50	•			
	Trp	Leu	Glu	Trp	Val	Lys	Gln	Arg	Pro	Gly	His	Gly	Leu	Glu	Trp	Val		
50			35	;				40)				45	i				

	Gly	Glu 50	Ile	Ser	Pro	Gly	Thr 55	Phe	Thr	Thr	Asn	Tyr 60	Asn	Glu	Lys	Phe	
5	Lys 65	Ala	Lys	Ala	Thr	Phe 70	Thr	Ala	Asp	Thr	Ser 75	Ser	Asn	Thr	Ala	Туг 80	
10	Leu	Gln	Leu	Ser	Gly 85	Leu	Thr	Ser	Glu	Asp 90	Ser	Ala	Val	Tyr	Phe 95	Cys	
15	Ala	Arg	Phe	Ser 100	His	Phe	Ser	Gly	Ser 105	Asn	Туr	Asp	Tyr	Phe 110	Asp	Туr	
20 ·	Trp	Gly	Gln 115	Gly	Thr	Ser	Leu	Thr 120	Val	Ser	Ser						
25	(2)	INF) SE		CE CI	HARA	CTER	isti(cs:	s							
30			(4	B) T C) S D) T	TRAN	DEDN	ESS:	dou									
35		(ii) MO:	LECU	LE T	YPE:	DNA	(ge	nomi	c)							
40		(ix	(A) N B) L	ame/ ocat	ION:	1	321	:/p	rodu	ct=	"lig	ht c	hain	var	iabl	e
45					do	ma i n	of	muri	ne a	ntib	ody	TES-	C21•				
50		(xi) SE	QUEN	CE D	ESCR	IPTI	CN:	SEQ	ID N	0:3:						

5	GAC	ATC	TTG	CTG	ACT	CAG	TY-T	CC4	ccc	יית ג	CITC	uv~m	~~~	B.Cm	001	001		40
			Leu															48
	1				5					10		-			15	GIY		
10	GAA	AGA	GTC	AGT	TTC	TCC	TGC	AGG	GCC	AGT	CAG	AGC	ATT	GGC	ACA	AAC		96
			Val															
				20					25					30				
15																		
			TGG															144
	116	His	Trp	Tyr	Gln	Gln	Arg		Asp	Gly	Ser	Pro			Leu	Ile		
			35					40					45	•				
20	AAG	TAT	GCT	••••••••••••••••••••••••••••••••••••	GAG	- TV-VIP	3.000	m~m	000	.								
			Ala															192
		50			014	251	55	Ser	GIA	116	PIO	60	Arg	Pne	Ser	GIY		
							-					ĢŪ						
25	AGT	GGA	TCA	GGG	ACA	GAG	TTT	ACT	CTA	AAC	ATC	AAC	AGT	GTG	GAG	TCT		240
			Ser															210
	65					70					75					80		
30																		
			ATT															288
	Glu	Asp	Ile	Ala	Asp	Tyr	Tyr	Cys	Gln	Gln	Ser	Asp	Ser	Trp	Pro	Thr		
					85					90					95	,		
35	.~~	~ ~	001	000														
			GGA									С						322
	1111	FIIE	Gly	100	GIY	unr	Lys	Leu		Ile	Lys							
				100					105									
40																		
	(2)	INFO	RMAT	ION	FOR	SEO	ID N	0:4:										
							•										•	
45		((i) S	EQUE	NCE	CHAR	ACTE	RIST	ics:									
					ngth					s								
			(B) TY	PE:	amin	o ac	id										

27

(D) TOPOLOGY: linear

50

		(ii	.) MC	DLECU	LE 1	YPE:	pro	tein	l							
5		(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:4:				•	
10	Asp 1		Leu	Leu	Thr 5		Ser	Pro	Ala	Ile 10	Leu	Ser	Val	Ser	Pro	Gly
•	Glu	Arg	Val	Ser 20	Phe	Ser	Cys	Arg	Ala 25	Ser	Gln	Ser	Ile	Gl <i>y</i> 30	Thr	Asn
15	Ile	His	Trp	Tyr	Gln	Gln	Arg	Thr 40	Asp	Gly	Ser	Pro	Arg 45	Leu	Leu	Ile
20	Lys	Tyr 50	Ala	Ser	Glu	Ser	Ile 55	Ser	Gly	Ile	Pro	Ser 60	Arg	Phe	Ser	Gly
25	Ser 65	Gly	Ser	Gly	Thr	Glu 70	Phe	Thr	Leu	Asn	11e 75	Asn	Ser	Val	Glu	Ser 80
30	Glu	Asp	Ile	Ala	Asp 85	Туr	Tyr	Cys	Ġĺn	Gln 90	Ser	Asp	Ser	Trp	Pro 95	Thr
35	Thr	Phe	Gly	Gly 100	Gly	Thr	Lys	Leu	Glu 105	Ile	Lys					
40	(2)		i	NOI												
		(i)	(A) TY () LE () TY	NGTH	1: 42	4 ba	se p	airs					-		•
4 5				c) si o) to					le							
50		(ii)	MOL	ECUL	Е ТҮ	PE:	DNA	(gen	omic)						

5		(i)	() FE	EATUF	Œ:													
	•		((A) N	IAME/	KEY:	CDS											
			(B) L	OCAT	'ION:	22.	.402					•				,	
10		:																
10		(ix	() FE															
			(A) N	IAME/	KEY:	mat	_pep	tide									
						: MOI												
15			(D) O	THER	INF	ORMA	TION	: /p	rodu	ct=	•lig	ht c	hain	var	iable		
13			•		re	gion	C21	-L1•										
				,					·									
20		(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:5:							
	CTC	CGCA	AGC	TTGC	CGCC	AC C											:	51
								Glu	Thr	Pro	Ala	Gln	Leu	Leu	Phe	Leu		
25 -							-20					-15						
	~~																•	
			CTG														9	99
			Leu	Trp	Leu		Asp	Thr	Thr	Gly	Asp	Ile	Leu	Leu	Thr	Gln		
30	-10					-5					1				5			
	300	200																
			GGC														14	17
	ser	Pro	Gly		Leu	Ser	Leu	Ser		Gly	Glu	Arg	Ala	Thr	Leu	Ser		
35				10					15					20				
	mcc.	3.00	000															
			GCC														19	15
	CYS	AIG	Ala	ser	GIN	ser	He		Thr	Asn	Ile	His		Tyr	Gln	Gln		
40			25		•			30	•				35					
	8.80	^^	000															
			GGC														24	13
	гåя		Gly	GIN	Alg	Pro		Leu	Leu	Ile	Lys	Tyr	Ala	Ser	Glu	Ser		
45		40					45	•				50						
	ייחל	100	~~~	3 ~~~	000													
			GGC														29	1
	55		Gly	TIE	PTO		Arg	Pne	Ser	Gly		Gly	Ser	Gly	Thr	_		
50	33					60					65					70		

29

. 55

5																	
															ATG		339
	Phe	Thr	Leu	Thr		Ser	Arg	Leu	Glu		Glu	Asp	Phe	Ala	Met	Tyr	•
					7 5					80					85		
10	~ 10		~~~			~1.0											
															GGC		387
	Tyr	Cys	GIN		ser	ASD	2er	тър		THE	Thr	Pne	GIY		Gly	The	
				- 90					95					100			
15		~~~		1 ma		0000	~ ~ ~ ~ ·		~~~			_			_	`	404
		GTG				CGIC	SAGT	ATT (CTAG	VAGG/	AT CO	3					424
	Lys	Val		116	rys												
			105														
20							_										
	(2)	T.177			200	200											
	(2)	INF	JKMA'	LION	FOR	SEQ	TD I	NO: 6	:								
			<i>1</i>	POIII	PMCE	CUAT	0 A ~m	CDTC	BICC								
25		,		_	ence Engti												
					YPE:				acro	15							
					OPOL												
			٠,	<i>)</i> , [(OFOLK		1111	SOL									
30		/225		. ECU	LE T	VDE.											
		(11)	, MOI	JECU1	DE 1.	IPE:	pro	rein									
		(vi	CF	TIEN!	CE D	בפרם.	T ውጥ T /	381. (ceo .	TD 37	٦						
35		(~ -	, 52,	20241	CD D	ooca.		J	JEQ .		J. U.						
33	Mor	Glu	Thr	Pro	פומ	Cln	Lou	Lou	Dho	1	1	1	7 ~		Leu	Desa	
	-20	GIU	1111	110	AIG	-15	Deu	LASU	riie	Leu	-10	reu	Leu	пр	ren	-5	
	20										-10					-3	
40	Asn	Thr	Thr	Gly	A STO	Tle	T.e.u	T.eu	Thr	Cln	Sor	Pro	Gly	ሞኮኮ	Leu	Sor	
	p			01,	1		Dea	Deu	5	GIII	Ser	110	GLY	10	Deu	261	
					•				,					10			
	ī.e.u	Ser	Pro	Glv	Glu	Δra	λla	Thr	Lau	50×	C/c	۸۳۵	212	cor	Gln	Cor	
45		001	15	013	010	g	****	20	200		Cys	AL 9	25	Ser	GIII	Ser	
			1,7					20					ده				
	Tle	Gly	ጥከታ	Δen	Tle	Hie	Tro	Tvr	Gla	Gla	Lve	Dro	Gl ₁ ,	Clr	۸ ۱-	Pro	
	***	30	****	പാവ	116		35	A y L		2111	шуз	40	ară	3111	WIG	FIO	
50		50					, ,					40					

	****			110	J	* y *	VIG	261	914	Ser	116	Ser	GIA	rie	FIO	Ser	
5	45					50			•		5 5					60	
	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Thr	Ile	Ser	
					65					70					75		
10	•																
	Arg	Leu	Glu	Pro	Glu	Asp	Phe	Ala	Met	Tyr	Tyr	Cys	Gln	Gln	Ser	Asp	
		•		80					85					90			-
15																	-
	Ser	Trp	Pro	Thr	Thr	Phe	Gly	Gln	Gly	Thr	Lys	Val	Glu	Ile	Lys		
	•		95					100					105				
20	(2)	INF	ORMA	PION	FOR	SEO	TD I	NO:7	•								
	,_,								•								•
		(i)) SE(OUEN	CE CI	HARAG	CTER	İSTI	cs:								
					ENGT					s							
25					YPE:												
			((c) s:	TRAN	DEDNI	ESS:	doul	ble								
			(1	D) TY	OPOL	OGY:	line	ear	,								
30																	
		(ii)) MOI	LECU	LE T	YPE:	DNA	(ge	nomi	c) .							
				·												•	
35								•									
••		(ix) FE	ATUR	E:												
			()	A) N	AME/I	KEY:	CDS										
			(1	B) Lo	OCAT:	ION:	22.	. 402									
40								•									
		(ix) FE	ATURI	E:												
			(1	A) N	AME/I	KEY:	mat_	pep	tide								
45					OCAT:												
•			(1	D) O	THER	INF	ORMA'	rion	: /p:	rodu	ct=	lig	ht c	hain	var	iable	j
					re	gion	C21	-L2°				÷					
50																	
~		(xi) SE(QUEN	CE DI	ESCR:	IPTI	ON: :	SEO :	ID N	0:7:						
								- '	•	•							

· 55

5																		
•	CIC	CGCA	AGC '	TTGC	CGCC	AC C	ATG	GAG	ACC	ccc	GCC	CAG	CTG	CTG	TTC	CTG	51	
							Met	Glu	Thr	Pro	Ala	Gln	Leu	Leu	Phe	Leu	•	
							-20					-15						
10	٠.																	
	CIG	CTG	CTG	TGG	CTG	ccc	GAC	ACC	ACC	GGC	GAC	ATC	CTG	CTG	ACC	CAG	99	
	Leu	Leu	Leu	Trp	Leu	Pro	Asp	Thr	Thr	Gly	Asp	Ile	Leu	Leu	Thr	Gln		
	-10					-5					1				5	•		
15		1																
	AGC	ccc	GGC	ACC	CTG	AGC	CTG	AGC	ccc	GGC	GAG	AGG	GCC	ACC	CTG	AGC	147	
	Ser	Pro	Gly	Thr	Leu	Ser	Leu	Ser	Pro	Gly	Glu	Arg	Ala	Thr	Leu	Ser		
				10		•			15					20				
20										•								
					CAG												195	
	Cys	Arg		Ser	Gln	Ser	Ile		Thr	Asn	Ile	His		Tyr	Gln	Gln		
			25					30					35					
25			222															
					GCC												243	
	ьуs		GIY	GIN	Ala	Pro		Leu	Leu	He	Lys		Ala	Ser	Glu	Ser		
••		40					45					50						
30	እ ጥ ና	A CC	ccc	N TVC	ccc	CAC	A C C	mm/c	NCC.	ccc	3.00	000	100	000	100	020	201	
					Pro												291	
	55	561	Gly	116	110	مور 60	MIG	FIIE	ser	GIY	65	GIÀ	Ser	GIY	TILL	70		
35						00					03					70		
	TTC	ACC	CTG	ACC	ATC	AGC	AGG	CTG	GAG	ccc	GAG	GAC	ጥጥ	CCC	ATY	ጥልር	339	
					Ile												333	
					75		3			80					85	-1-		
40											•							
	TAC	TGC	CAG	CAG	AGC	GAC	AGC	TGG	ccc	ACC	ACC	TTC	GGC	CAG	GGC	ACC	387	
					Ser													
				90		, -			95					100				
45									_									
•	AAG	GTG	GAG	ATC	AAA	CGTY	GAGT	ATT (CTAG	AAGG	AT C	3					424	
	Lys	Val	Glu	Ile	Lys											•		
			105															

5	(2)	INF	ORMA	TION	FOR	SEQ	ΙŅΙ	8 : OM	:							
10	·. 	•	C	SEQU A) L B) T	ENGT YPE:	H: 1 ami	27 au no a	mino cid								
15) MO						SEQ :	ID N	D:8:					
20	Met -20	Glu	Thr	Pro	Ala	Gln -15	Leu	Leu	Phe	Leu	Leu -10	Leu	Leu	Trp	Leu	Pro -5
25	Asp	Thr	Thr	Gly	Asp 1	Ile	Leu	Leu	Thr 5	Gln	Ser	Pro	Gly	Thr 10	Leu	Ser
30 '			15					20					25		Gln	
35		30					35					40		į	λla	
	45					50					55				Pro	60
4 0					. 65					70					11e 75	
45				80					85		,			90	Ser	Asp
50	ser	ттħ	Pro 95	inr	Thr	rne	GIÀ	100	Gly	Thr	Lys	Val	Glu 105	Ile	Lys	

5	(2) INFORMATION FOR SEQ) ID NO:9:	
10	(B) TYPE: nuc	24 base pairs cleic acid	
	(C) STRANDEDN (D) TOPOLOGY:		
15	(ii) MOLECULE TYPE:	DNA (genomic)	
٠			
20	(ix) FEATURE:		
	(A) NAME/KEY:	CDS	
	(B) LOCATION:	22402	
25	(ix) FEATURE:		
	(A) NAME/KEY:	mat_peptide	
	(B) LOCATION:		
	, , , , , , , , , , , , , , , , , , , ,		
		·	
30	(D) OTHER INF	ORMATION: /product= "light chain variable C21-L3"	
30	(D) OTHER INF	ORMATION: /product= *light chain variable	
30	(D) OTHER INF region	ORMATION: /product= *light chain variable C21-L3*	
30	(D) OTHER INF region	ORMATION: /product= *light chain variable	
	(D) OTHER INF region (xi) SEQUENCE DESCR	ORMATION: /product= *light chain variable C21-L3*	
	(D) OTHER INF region (xi) SEQUENCE DESCR	ORMATION: /product= *light chain variable . C21-L3* IPTION: SEQ ID NO:9: ATG GAG ACC CCC GCC CAG CTG CTG TTC CTG 51	
	(D) OTHER INF region (xi) SEQUENCE DESCR	ORMATION: /product= *light chain variable C21-L3*	
	(D) OTHER INF region (xi) SEQUENCE DESCR	ORMATION: /product= *light chain variable C21-L3* IPTION: SEQ ID NO:9: ATG GAG ACC CCC GCC CAG CTG CTG TTC CTG 51 Met Glu Thr Pro Ala Gln Leu Leu Phe Leu	
35	(D) OTHER INF region (xi) SEQUENCE DESCR CTCCGCAAGC TTGCCGCCAC C	ORMATION: /product= *light chain variable C21-L3* IPTION: SEQ ID NO:9: ATG GAG ACC CCC GCC CAG CTG CTG TTC CTG 51 Met Glu Thr Pro Ala Gln Leu Leu Phe Leu	
35	(D) OTHER INF region (xi) SEQUENCE DESCR CTCCGCAAGC TTGCCGCCAC C CTG CTG CTG TGG CTG CCC	ORMATION: /product= *light chain variable C21-L3* IPTION: SEQ ID NO:9: ATG GAG ACC CCC GCC CAG CTG CTG TTC CTG Met Glu Thr Pro Ala Gln Leu Leu Phe Leu -20 -15	
35	(D) OTHER INF region (xi) SEQUENCE DESCR CTCCGCAAGC TTGCCGCCAC C CTG CTG CTG TGG CTG CCC	ORMATION: /product= *light chain variable C21-L3* IPTION: SEQ ID NO:9: ATG GAG ACC CCC GCC CAG CTG CTG TTC CTG 51 Met Glu Thr Pro Ala Gln Leu Leu Phe Leu -20 -15 GAC ACC ACC GGC GAG ATC GTG CTG ACC CAG 99 Asp Thr Thr Gly Glu Ile Val Leu Thr Gln	
35 40	(D) OTHER INF region (xi) SEQUENCE DESCR CTCCGCAAGC TTGCCGCCAC C CTG CTG CTG TGG CTG CCC Leu Leu Leu Trp Leu Pro -10 -5	ORMATION: /product= *light chain variable C21-L3* IPTION: SEQ ID NO:9: ATG GAG ACC CCC GCC CAG CTG CTG TTC CTG 51 Met Glu Thr Pro Ala Gln Leu Leu Phe Leu -20 -15 GAC ACC ACC GGC GAG ATC GTG CTG ACC CAG 99 Asp Thr Thr Gly Glu Ile Val Leu Thr Gln	
35 40	(D) OTHER INF region (xi) SEQUENCE DESCR CTCCGCAAGC TTGCCGCCAC C CTG CTG CTG TGG CTG CCC Leu Leu Leu Trp Leu Pro -10 -5 AGC CCC GGC ACC CTG AGC	ORMATION: /product= *light chain variable C21-L3* IPTION: SEQ ID NO:9: ATG GAG ACC CCC GCC CAG CTG CTG TTC CTG 51 Met Glu Thr Pro Ala Gln Leu Leu Phe Leu -20 -15 GAC ACC ACC GGC GAG ATC GTG CTG ACC CAG 99 Asp Thr Thr Gly Glu Ile Val Leu Thr Gln 1 5	

3																		
	TGC	AGG	GCC	AGC	CAG	AGC	ATC	GGC	ACC	AAC	ATC	CAC	TGG	TAC	CAG	CAG		195
	Cys	Arg	Ala	Ser	Gln	Ser	Ile	Gly	Thr	Asn	Ile	His	Trp	Tyr	Gln	Gln		
10			25					30					35					
,,,																		
	AAG	ccc	GGC	CAG	GCC	ccc	AGG	CTG	CTG	ATC	AAG	TAC	GCC	AGC	GAG	AGC		243
	Lys	Pro	Gly	Gln	Ala	bio	Arg	Leu	Leu	Ile	Lys	Tyr	Ala	Ser	Glu	Ser	•	
15		40					45					50						
		AGC																291
		Ser	Gly	Ile	Pro	Ser	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp		
20	55					60					65	2*	. •			70		
•		ACC													,			339
	Phe	Thr	Leu	Thr		Ser	Arg	Leu	Glu		Glu	Asp	Phe	Ala		Tyr		
25					75					80					85			
	m >0	5 00	222															
•		TGC																387
	туг	Cys	GIN		Ser	Asp	Ser	Trp		Thr	Thr	Phe	Gly		Gly	Thr		
30				90					95					100			•	
	AAC	GTG	CAG	λTV		CON	~ A CTT:	a mora	~!!		. m c	,						424
		Val				CGI	3AG:17	,,,	LING	MGG	AT C	-						424
	2,5	vui	105	116	Llys													
35			103															
	(2)	INFO	ORMAT	CION	FOR	SEO	ID 1	W:10) <u></u>									
										•								
ю			(i) 5	SEQUI	ENCE	CHA	RACTI	ERIS	rics	:								
					ENGT													
					YPE:													
, (5			(I) T (OPOLO	ŒΥ:	line	ear										
		(ii)	HOI	ECUI	LE TY	PE:	prot	.ein										
ю		(xi)	SEC	QUENC	CE DI	SCR	IPTI	N: :	SEQ :	ID N	0:10	:						

. 5	Met -20	Glu	Thr	Pro	Ala	Gln -15	Leu	Leu	Phe	Leu	Leu -10	Leu	Leu	Trp	Leu	Pro -5
10	Asp	Thr	Thr	Gly	Glu 1	Ile	Val	Leu	Thr 5	Gln	Ser	Pro	Gly	Thr 10	Leu	Ser
15	Leu	Ser	Pro 15	Gly	Glu	Arg	Ala	Thr 20	Leu	Ser	Cys	Arg	Ala 25	Ser	Gln	Ser
	Ile	Gly 30	Thr	Asn	Ile	His	Trp 35	Tyr	Gln	Gln	Lys	Pro 40	Gly	Gln	Ala	Pro
20	Arg 45	Leu	Leu	Ile	Lys	Tyr 50	Ala	Ser	Glu	Ser	Ile 55	Ser	Gly	Ile	Pro	Ser 60
25	Arg	Phe	Ser	Gly	Ser 65	Gly	Ser	Gly	Thr	Asp 70	Phe	Thr	Leu	Thr	Ile 75	Ser
30	Arg	Leu	Glu	Pro 80	Glu	Asp	Phe	Ala	Met 85	Туr	Туr	Cys	Gln	Gln 90	Ser	Asp
35	Ser	Trp	Pro 95	Thr	Thr	Phe	Gly	Gln 100	Gly	Thr	Lys	Val	Glu 105	Ile	Lys	
	(2)	INFO	ORMAI	MOIT	FOR	SEQ	ID 1	10:11	l:				٠			
40		(i)	SEC	-	CE CH					5						
45 -			(0	c) si	(PE: TRANI OPOLO	DEDNE	ss:	doul		,						
50		(ii)	MOI	LECUI	E T	PE:	DNA	(ger	nomic	:)						

5		(ix)	FEA	TURE	:												
			(2	A) NA	ME/F	EY:	CDS										
	٠.		{ E	3) LC	CATI	ON:	22	447									
10		(ix)	FEA	TURE	::			•									
			{2	A) NA	ME/I	EY:	mat_	pept	ide								
			(E	3) LC	CAT	ON:	79	447									
			(1) Q 1	HER	INFO	ORMAT	PION:	/pı	coduc	:t= '	'hea v	vy c)	nain	vari	iable	
15					reg	jion	C21-	-н1 •					•				
		•															
20		(xi)	SEC	QUENC	E DI	ESCR	[PTIC	ON: S	SEQ 1	ED NO):11:	:					
	CTC	CGCAZ	AGC 1	MGCC	GCC	AC C	ATG	GAC	TGG	ACC	TGG	AGG	GTG	TTC	TGC	CTG	51
							Met	Asp	Trp	Thr	Trp	Arg	Val	Phe	Cys	Leu	•
25			٠.		,		-19				-15					-10	
					,												
	CTG	GCC	GTG	GCC	CCC	GGC	GCC	CAC	AGC	CAG	GTG	CAG	CTG	GTG	CAG	AGC	99
	Leu	Ala	Val	Ala	Pro	Gly	Ala	His	Ser	Gln	Val	Gln	Leu	Val	Gln	Ser	
30					-5					1				5			
	GGC	GCC.	GAG	GTG	AAG	AAG	ccc	GGC	GCC	AGC	GTG	AAG	GTG	AGC	TGC	AAG	147
	Gly	Ala		Val	Lys	Lys	Pro	_	Ala	Ser	Val	Lys		Ser	Cys	Lys	
35			. 10					15					20				
		AGC				•											195
	Ala	Ser	Gly	Tyr	Thr	Phe		Met	Tyr	Trp	Leu		Trp	Val	Lys	Gln	
40		25					30					35					
	AGG	CCC	GGC	CAC	GGC	CTG	GAG	TGG	GTG	GGC.	GAG	ATC	AGC	ccc	GGC	ACC	243
45	Arg	Pro	Gly	His	Gly	Leu	Glu	Trp	Val	Gly	Glu	Ile	Ser	Pro	Gly	Thr	
40	40					45					50					55	
												•					
		ACC															291
50	Phe	Thr	Thr	Asn	Tyr	Asn	Glu	Lys	Phe	Lys	Ala	Lys	Ala	Thr	Phe	Thr	
					60					65					70		

37

. 55

5													•			•	
	GCC	GAC	ACC	AGC	ACC	AAC	ACC	GCC	TAC	ATG	GAG	CTG	AGC	AGC	CTG	ACC	339
	Ala	Asp	Thr	Ser	Thr	Asn	Thr	Ala	Туr	Met	Glu	Leu	Ser	Ser	Leu	Thr	
				75					80					85			
10																	
	AGC	GAG	GAC	ACC	GCC	GTG	TAC	TAC	TGC	GCC	AGG	TTC	AGC	CAC	TTC	AGC	387
	Ser	Glu	Asp	Thr	Ala	Val	Tyr	Tyr	Cys	Ala	Arg	Phe	Ser	His	Phe	Ser	•
			90					95					100				
15																•	
	GGC	AGC	AAC	TAC	GAC	TAC	TTC	GAC	TAC	TGG	GGC	CAG	GGC	ACC	CTG	GTG	435
	Gly	Ser	Asn	Tyr	Asp	Tyr	Phe	Asp	Tyr	Trp	Gly	Gln	Gly	Thr	Leu	Va1	
		105					110					115				•	
20				•													
	ACC	GTG	AGC	TCA	GGT	GAGT	ICT I	AGAA	GGGA'	rc c							468
	Thr	Val	Ser	Ser													
	120																
25																	
	(2)	INF	ORMA'	TION	FOR	SEQ	ID I	NO:1	2:								
00										-							
30			(i)	SEQU	ENCE	CHA	RACT	ERIS	TICS	:							4
			(,	A) L	ENGT	H: 1	42 aı	mino	aci	ds							
			C	B) T	YPE:	ami	no a	cid									
35			C	D) T	OPOL	OGY:	lin	ear									
33																	
		(ii) MO	LECU:	LE T	YPE:	pro	tein				•					
40		(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:12	:					
••																	
	Met	Asp	Trp	Thr	Trp	Arg	Val	Phe	Cys	Leu	Leu	Ala	Val	Ala	Pro	Gly	
•	-19				-15					-10			•		-5	÷	
45																	
	Ala	His	Ser	Gln	Val	Gln	Leu	Val	Gln	Ser	Gly	Ala	Glu	Val	Lys	Lys	
				1				5			_		10		-		

_	Pro	Gly 15	Ala	Ser	Val	Lys	Va1	Ser	Cys	Lys	Ala	Ser 25	Gly	Туг	Thr	Phe	
<i>5</i>		Met	Tyr	Trp	Leu		Trp	Val	Lys	Gln	-	Pro	Gly	His	Gly		
10	30 Glu	Trp	Val	Gly	Glu	35	Ser	Pro	Gly	Thr	40 Phe	Thr	Thr	Asn	Tyr	45 Asn	
					50					55				•	60		
	Glu	Lys	Phe	Lys 65	Ala	Lys	Ala	Thr	Phe 70	Thr	Ala	Asp	Thr	Ser - 75	Thr	Asn	
20	Thr	Ala	T yr 80	Met	Glu	Leu	Ser	Ser 85	Leu	Thr	Ser	Glu	Asp 90	Thr	Ala	Val	
25	Туr	Tyr 95	Cys	Ala	Arg	Phe	Ser 100	His	Phe	Ser	Gly	Ser 105	Asn	Tyr	Asp	Tyr	
30	Phe 110	Asp	Tyr	Trp	Gly	Gln 115	Gly	Thr	Leu	Val	Thr 120		Ser	Ser			
	(2)	INF	ORMA'	rion	FOR	SEQ	ID	NO:1	3:								
35 ·		(i	· (A) L	CE C	H: 4	68 b	ase :	pair	s							
40			(c) s	YPE: TRAN OPOL	DEDN	ESS:	dou									
45		(ii) MO:	LECU	LE T	YPE:	DNA	(ge	nomi	c)							
		(ix) FE.				ana.										
50		•			ame/ ocat												

5		(ix)	FEA	ATURI	::						-						
			(2	() N	ME/	ŒY:	mat_	pept	ide								•
			(E	3) L(CAT	ON:	79.	447									
10			(I	o) 01	MER	INFO	RMAT	'ION	/pı	coduc	:t= '	heav	y ch	nain	vari	able	
					reg	jion	C21-	-н3•									
				•													
15		(xi)	SEC	DUENC	E DE	ESCRI	PTIC	XV: 5	SEQ I	ID N	0:13:	:					
	CICC	:GCA	AGC 1	rtgcc	.GCC2	AC C											51
,								ASP	TŢ	Thr	_	Arg	Vai	Pne	Cys		
20							-19				-15			-		-10	
	C IV	ccc	CTY.	GCC	ccc	ccc	ccc	CNC	ACC.	CNG	CTC	CNC	CTC	CINC	CAG	AGC.	99
				Ala													
25	Den	AIG	·	ALG	-5	Gry	AIG	*****	361	1	401	GIII	Deu	5		Jer	
			•		•									•			
	GGC	GCC	GAG	GTG	AAG	AAG	ccc	GGC	GCC	AGC	GTG	AAG	GTG	AGC	TGC	AAG	147
	Gly	Ala	Glu	Val	Lys	Lys	Pro	Gly	Ala	ser	Val	Lys	Val	Ser	Cys	Lys	
30			10					15					20				
													,				
	GCC	AGC	GGC	TAC	ACC	TTC	AGC	ATG	TAC	TGG	CTG	GAG	TGG	GTG	AGG	CAG	195
35	Ala		Gly	Tyr	Thr	Phe		Met	Tyr	Trp	Leu	Glu	Trp	Val	Arg	Gln	
33		25					30					35					
														•			
				CAC													243
40		Pro	Gly	His	Gly		Glu	Trp	Val	Gly		Ile	Ser	Pro	Gly		
	40					45					50					55	
	-																201
																ACC	291
45	Pne	Tnr	Thr	Asn	-	ASN	GIU	Lys	Pne	-	A19	Arg	Ala	Thr		TNY	
					60					. 65					70		

									-								
5	GCC	GAC	ACC	AGC	ACC	AAC	ACC	GCC	TAC	ATG	GAG	CTG	AGC	AGC	CIG	AGG	339
	Ala	Asp	Thr	Ser	Thr	Asn	Thr	Ala	Tyr	Met	Glu	Leu	Ser	Ser	Leu	Arg	
				75					80					85			
10 .																	
	AGC	GAG	GAC	ACC	GCC	GTG	TAC	TAC	TGC	GCC	AGĠ	TTC	AGC	CAC	TTC	AGC	387
	Ser	Glu	Asp	Thr	Ala	Val	Туr	Tyr	Cys	Ala	Arg	Phe	Ser	His	Phe	Ser	
			90					95					100				
15																	
	GGC	AGC	AAC	TAC	GAC	TAC	TTC	GAC	TAC	TGG	GGC	CAG	GGC	ACC	CTG	GTG	435
	Gly	Ser	Asn	Tyr	Asp	Tyr	Phe	Asp	Tyr	Trp	Gly	Gln	Gly	Thr	Leu	Val	
		105					110					115					
20															•		
			AGC		GGT	GAGT	ICT I	AGAA	GGGA!	rc c							468
		Val	Ser	Ser													
	120																
25			•														
	121	TAIC	ORMA!	PT (NI	EOB	CEO	TD :	10.1	4.							•	
	(2)	THE	JRPA.	11011	FOR	SEQ	10	WO. I	*:		•					•	
30			(i) :	SEOU	ENCE	CHA	RACT	ERIS	TICS	:							
30				A) L													
•			(1	B) T	YPE:	ami	no a	cid									
		٠	{ 1	D) Tr	OPOL	OGY;	lin	ear								•	
35																•	
		(ii) MO	LECU	LE T	YPE:	pro	tein								•	
								•									
•		(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:14	:					
40																	
	Met	Asp	Trp	Thr	Trp	Arg	Val	Phe	Cys	Leu	Leu	Ala	Val	Ala	Pro	Gly	
	-19				-15	-				-10					-5		
45	Ala	His	Ser	Gln	Val	Gln	Leu	Val	Gln	Ser	Gly	Ala	Glu	Val	Lys	Lys	
	٠.			1				5					10				

5	Pro	Gly 15	Ala	Ser	Val	Lys	Va1 20	Ser	Cys	Lys	Ala	Ser 25	Gly	Tyr	Thr	Phe
10	Ser 30	Met	Туr	Trp	Leu	Glu 35	Trp	Val	Arg	Gln	Ala 40	Pro	Gly	His	Gly	Leu 45
15	Glu	Trp	Val	Gly	Glu 50	Ile	Ser	Pro	Gly	Thr 55	Phe	Thr	Thr	Asn	Tyr 60	Asn
	Glu	Lys	Phe	Lys 65	λla	λrg	Ala	Thr	Phe 70	Thr	Ala	Asp	Thr	Ser 75	Thr	Asn
20	Thr	Ala	Tyr 80	Met	Glu	Leu	Ser	Ser 85	Leu	Arg	Ser	Glu	Asp 90	Thr	Ala	Val
25	Tyr	Tyr 95	Cys	Ala	Arg	Phe	Ser 100	His	Phe	Ser	Gly	Ser 105	Asn	Tyr	Asp	Tyr
30	Phe 110	Asp	Tyr	Trp	Gly	Gln 115	Gly	Thr	Leu	Val	Thr 120	Val	Ser	Ser		
35	(2)	INF	ORMA			,										
40		,	(1	A) L1 B) T' C) S' D) T(YPE: TRAN	nuc DEDN	leic ESS:	acio	d	S.						
45		(ii) MO	LECU	LE T	YPE:	DNA	(ge	nomi	c)			l,	٠	1	
50		(ix		ATUR A) N B) L	ame/			•								

5		(ix	FE.	ATURI	E:												
			(2	A) N/	AME/I	KEY:	mat.	pep	tide								•
			(1	B) LO	CAT:	ON:	79.	.447									
			(1	o) o:	THER	INF	ORMA	rion.	: /p:	rodu	ct=	hea '	vy cl	nain	var	iable	
10					re	gion	C21	-llay	1•								
					•			•									
	•	(xi)	SE	QUEN(CE D	ESCR:	IPTI	:: MC	SEQ :	ID N	0:15	:					
15																	
	CTC	CGCA	AGC :	rtgc	CGCC	AC C	ATG	GAC	TGG	ACC	TGG	AGG	GTG	TTC	TGC	CTG	51
		-					Met	Asp	Trp	Thr	Trp	Arg	Val	Phe	Cys	Leu	
							-19				~15					-10	
20																	
				GCC													99
	Leu	Ala	Val	Ala		Gly	Ala	His	Ser		Val	Gln	Leu		Gln	Ser	
25					-5					. 1				5			
	ccc	CCC	~~	~~			000		000				~~~				
				Val												AAG	147
	GIY	WIG	10	Val	Lys	Lys	PIO	15	Ala	ser	vai	Lys	20	Ser	Cys	Lys	
30			10					13					20				
	GCC	AGC	GGC	TAC	ACC	77 C	AGC	ATY	ጥልር	TCC	CTC	GAG	TGG	GTG	AAC	CAG	195
				Tyr													1,,,
		25	2	-, -			30		-1-			35	LLP		D , 0	0111	
35																	
	AGG	ccc	GGC	CAG	AGG	CTG	GAG	TGG	ATG	GGC	GAG	ATC	AGC	ccc	GGC	ACC	243
				Gln													
	40					45				_	50				-	55	
40																	
	TTC	ACC	ACC	AAC	TAC	AAC	GAG	AAG	TTC	AAG	GCC	AAG	GCC	ACC	TTC	ACC	291
	Phe	Thr	Thr	Asn	Tyr	Asn	Glu	Lys	Phe	Lys	Ala	Lys	Ala	Thr	Phe	Thr	
45	,				60					65					70		

5																	
	GCC	GAC	ACC	AGC	GCC	AGC	ACC	GCC	TAC	ATG	GAG	CTG	AGC	AGC	CTG	ACC	339
	Ala	Asp	Thr	Ser	Ala	Ser	Thr	Ala	Tyr	Met	Glu	Leu	Ser	Ser	Leu	Thr	
				7Š					80					85			
10																	
	AGC	GAG	GAC	ACC	GCC	GTG	TAC	TAC	TGC	GCC	AGG	TTC	AGC	CAC	TTC	AGC	387
	Ser	Glu	Asp	Thr	Ala	Val	Tyr	Tyr	Сув	λla	Arg	Phe	Ser	His	Phe	Ser	
			90					95					100				
15																	
			AAC														435
	Gly		Asn	Tyr	Asp	Tyr	Phe	Asp	Tyr	Trp	Gly	Gln	Gly	Thr	Leu	Val	
		105					110					115					
20																	
			AGC		GGT	GAGT	rct i	AGAA	GATY	œ							467
		Val	Ser	Ser			•										
	120		•													•	
25				•									•				
	(2)	TNF	ORMA!	rton	FOR	SEO	TD (راد مرار ا	ς.								
	,				,	424											
			(i) :	SEQUI	ENCE	CHAI	RACT	ERIS	rics	•							
30			(1	A) L	engti	H: 14	12 ar	nino	acio	is							
			, a	B) T	YPE:	amin	no a	cid									
			. (1	D) T(OPOL	ŒΥ:	line	ear									
35		(ii	MOI	LECUI	LE T	YPE:	pro	ein									
		(xi	SE(QUEN	CE DI	ESCR:	[PTI	ON: :	SEQ I	ID NO	0:16	:					
									•						,		
40	Met	Asp	Trp	Thr	Trp	Arg	Val	Phe	Cys	Leu	Leu	Ala	Val	Ala	Pro	Gly	
	-19				-15					-10					-5		
		-															
	Ala	His	Ser	Gln	Val	Gln	Leu	Val	Gln	Ser	Gly	Ala	Glu	Val	Lys	Lys	
45				1				5					10				
								•									

5	Pro	Gly 15	Ala	Ser	Val.	Lys	Val 20	Ser	Cys	Lys	Ala	Ser 25	Gly	Tyr	Thr	Phe
10	Ser 30	Met	Туr	Trp	Leu	Glu 35		Val	Lys	Gln	Arg 40	Pro	Gly	Gln	Arg	Leu 45
15	Glu	Trp	Met	Gly	Glu 50	Ile	Ser	Pro	Gly	Thr 55	Phe	Thr	Thr	Asn	Туг 60	Asn
20	Glu	Lys	Phe	Lys 65	Ala	Lys	Ala	Thr	Phe 70	Thr	Ala	Asp	Thr	Ser 75	Ala	Ser
	Thr	Ala	Туг 80	Met	Glu	Leu	Ser	Ser 85	Leu	Thr	Ser	Glu	A sp 90	Thr	λla	Val
25	Tyr	Tyr 95	Cys	Ala	Arg	Phe	Ser 100	His	Phe	Ser	Gly	Ser 105	Asn	Tyr	λsp	Tyr
30	Phe 110	Asp	Tyr	Trp	Gly	Gln 115	Gly	Thr	Leu	Val	Thr 120	Val	Ser	Ser		
35	(2)						ID I						·			
40		(1)	(1	A) LI B) T	ENGTI YPE: TRANI	nuc DEDN	CTER: 67 ba leic ESS: line	ase p acid	pair:	s						
4 5		(ii					DNA		nomi	c)						
50		(ix	(A) N		KEY:	, CDS 22.	. 447								
55																

		(ix)	FEA	TURE	S:									•				
			(A	A) NA	ME/K	EY:	mat_	pept	ide									
			(E	3) LC	CATI	ON:	79	447										
10			(1	o) O1	HER	INFO	RMAT	NOI!	/pr	ođuc	t= •	heav	y ch	ain	vari	able		
					reg	ion	C21-	Науз	3 =									
	•	*																
15																		
		(xi)	SE(QUENC	CE DE	SCRI	PTI	: 1K	SEQ 1	ID NO	:17:	:						
20	CTCC	GCA	AGC 7	rrgco	CGCC#	AC C	ATG	GAC	TGG	ACC	TGG	AGG	GTG	TTC	TGC	CTG	51	
20							Met	Asp	Trp	Thr	Trp	Arg	Val	Phe	Cys	Leu		
							-19				-15					-10		
		·																
25	CTG	GCC	GTG	GCC	ccc	GGC	GCC	CAC	AGC	CAG	GTG	CAG	CTG	GTG	CAG	AGC	99	
20	Leu	Ala	Val	Ala	Pro	Ġly	Ala	His	Ser	Gln	Val	Gln	Leu	Val	Gln	Ser		
					-5					1		:		5				
30				GTG													147	
	Gly	Ala		Val	Lys	Lys	Pro	_	Ala	Ser	Val	Lys		Ser	Cys	Lys		
			10					15					20					
			~~~	<b></b>		-			<b></b> .	maa	000	a. a	maa	omo.		010	195	
35				TAC												_	193	
	Ala		-	Tyr	Thr	Pne		Met	Tyr	Trp	Leu		Trp	vaı	Arg	Gin		
		25					30					35						
	ccc	ccc	ccc	CAG	A G G	CTC	GAG	TYCC	ልጥር	ccc	GAG	ልጥዮ	እርር	ccc	ccc	ACC	243	
40																Thr		
	40	PLO	GIY	GIII	AIG.	45	GIU	110	Hec	GIY	50		Ser	110	Gry	55		
	₩.0	•				40					J.					در		
	<b>JATA</b> C	) ACC	ልቦር	220	ጥልር	<b>A</b> A C	GAG	V V C	ባግሃጉ	AAC:	GCC	ልርር		204	<b>ም</b> ጥ	ACC	291	
45																Thr		
÷	* 116			non	60		O.Lu	د رب		65		urg			70			
										0.5								

5																		
	CCC	GAC	ACC	AGC	GCC	AGC	ACC	GCC	TAC	ATG	GAG	CTG	AGC	AGC	CTG	AGG		339
	Ala	qaA	Thr	Ser	Ala	Ser	Thr	Ala	Tyr	Met	Glu	Leu	Ser	Ser	Leu	Arg		
				75					80					85				
10																		
	AGC	GAG	GAC	ACC	GCC	GTG	TAC	TAC	TGC	GCC	AGG	TTC	AGC	CAC	TTC	AGC		387
	Ser	Glu	Asp	Thr	Ala	Val	Tyr	Tyr	Cys	Ala	Arg	Phe	Ser	His	Phe	Ser		
			90					95					100					
15						-												
	GGC	AGC	AAC	TAC	GAC	TAC	TTC	GAC	TAC	TGG	GCC	CAG	GGC	ACC	CTG	GTG		435
	Gly	Ser	Asn	Tyr	Asp	Tyr	Phe	Asp	Tyr	Trp	Gly	Gln	Gly	Thr	Leu	Val		
		105				-	110	•				115						
20																	,	
£	ACC	GTG	AGC	TCA	GGTY	SAGT	PCT A	AGAA	GGATY	œ								467
	Thr	Val	Ser	Ser														
	120																	
25																		
		•																
	(2)	INF	ORMA	rion	FOR	SEQ	ID I	NO:1	B:									
30	٠		(i) :	SEQU	ENCE	CHA	RACTI	ERIS	TICS	:				,				
			C	A) L	ENGTI	H: 1	42 ar	nino	aci	ds								
			(1	B) T	YPE:	amiı	no ac	cid										
			(1	D) T	OPOL	OGY:	line	ear			-							
35																		
		(ii	) MO	LECU	LE T	YPE:	prot	tein										
		•																
		(xi	) SE	QUEN	CE D	ESCR:	IPTI	ON: :	SEQ :	ID N	0:18	:						
40																		
	Met	Asp	Trp	Thr	Trp	Arg	Val	Phe	Cys	Leu	Leu	Ala	Val	Ala	Pro	Gly		
	-19				-15					-10		•			-5			
	•																	
45	Ala	His	Ser	Gln	Val	Gln	Leu	Val	Gln	Ser	Gly	Ala	Glu	Val	Lys	Lys		
				1				5					10					

<b>.</b>	Pro	Gly 15	Ala	Ser	Val	Lys	Val . 20	Ser	Cys	Lys	Ala	Ser 25	Gly	Tyr	Thr	Phe
10	Ser 30	Met	Туг	Trp	Leu	Glu 35	Trp	Val	Arg	Gln	Ala 40	Pro	Gly	Gln	Arg	Leu 45
15	Glu	Trp	Met	Gly	Glu 50	Ile	Ser	Pro	Gly	Thr 55	Phe	Thr	Thr	Asn	Tyr 60	Asn
	Glu	Lys	Phe	Lys 65	Ala	Arg	Ala	Thr	Phe 70	Thr	Ala	Asp	Thr	Ser 75	Ala	Ser
20	Thr	Ala	Туг 80	Met	Glu	Leu	Ser	Ser 85	Leu	Arg	Ser	Glu	Asp 90	Thr	Ala	Val
25	Tyr	<b>Tyr</b> 95	Cys	Ala	Arg	Phe	Ser 100	His	Phe	Ser	Gly	Ser 105	Asn	Tyr	Asp	Tyr
30	Phe	Asp	Tyr	Trp	Gly	Gln 115	Gly	Thr	Leu	Val	Thr 120	Val	Ser	Ser		
35	(2)			TION												
40		. (1)	() ()	QUENCA) LI B) T	engti Ype:	nuc	05 ba	ase p	pair: d	S						
			(1	C) S'	OPOL	ÖGY:	line	ear	_							
45		(ii)	) MO	LECU	LE T	YPE:	DNA	(ge:	nomi	c)						
50		(xi)	) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:19	:				

6	TGAAGAAAGC TTGCCGCCAC CATGGAGACC CCCGCCCAGC TGCTGTTCCT GCTGCTGCTG	. 60
	TGGCTGCCCG ACACCACCGG CGACATCCTG CTGACCCAGA GCCCC	105
10	(2) INFORMATION FOR SEQ ID NO:20:	
	(1) SEQUENCE CHARACTERISTICS:	
15	(A) LENGTH: 95 base pairs	
••	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
20		
	(ii) MOLECULE TYPE: DNA (genomic)	
-		
25	(with CDOURNOR BECONTONION and The No. of	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:	
	GATGTTGGTG CCGATGCTCT GGCTGGCCCT GCAGCTCAGG GTGGCCCTCT CGCCGGGGCT	60
30		•
	CAGGCTCAGG GTGCCGGGGC TCTGGGTCAG CAGGA	95
		·
	(2) INFORMATION FOR SEQ ID NO:21:	
35		
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 75 base pairs	
	(B) TYPE: nucleic acid	
40	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(44) NOT FOUND THE ASSESSMENT OF THE ASSESSMENT	
45	(ii) MOLECULE TYPE: DNA (genomic)	
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	-

5	CAGAGCATCG GCACCAACAT CCACTGGTAC CAGCAGAAGC CCGGCCAGGC CCCCAGGCTG	60
	CTGATCAAGT ACGCC	75
10	(2) INFORMATION FOR SEQ ID NO:22:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 87 base pairs	
15	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
20	(ii) MOLECULE TYPE: DNA (genomic)	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	
	AGGGTGAAGT CGGTGCCGCT GCCGCTGCCG CTGAACCTGC TGGGGATGCC GCTGATGCTC	. 60
30	TCGCTGGCGT ACTTGATCAG CAGCCTG	87
	(2) INFORMATION FOR SEQ ID NO:23:	
35	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 84 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
40	(D) TOPOLOGY: linear	
٠	(ii) MOLECULE TYPE: DNA (genomic)	
45		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:	

	GCGGCACCGA CTTCACCCTG ACCATCAGCA GGCTGGAGCC CGAGGACTTC GCCATGTACT	60
	ACTGCCAGCA GAGCGACAGC TGGC	- 84
10	(2) INFORMATION FOR SEQ ID NO:24:	
	(i) SEQUENCE CHARACTERISTICS:	
15	(A) LENGTH: 82 base pairs	
13	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
20		
	(ii) MOLECULE TYPE: DNA (genomic)	
25		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	*,
	TTIGGATECT TETAGAATAE TEACGITTGA TETECACETT GGTGCCETGG CEGAAGGTGG	. 60
30		
	TGGGCCAGCT GTCGCTCTGC TG	82
	(2) INFORMATION FOR SEQ ID NO:25:	
35		
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 97 base pairs	
	(B) TYPE: nucleic acid	
40	(C) STRANDEDNESS: single	•
	(D) TOPOLOGY: linear	
45	(ii) MOLECULE TYPE: DNA (genomic)	
	(vi) CECHENCE DECORIDATION, CEO ID NO.25.	
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	

<b>.</b>	TGAAGAAAGC TTGCCGCCAC CATGGACTGG ACCTGGAGGG TGTTCTGCCT GCTGGCCGTG	60
	GCCCCCGGCG CCCACAGCCA GGTGCAGCTG GTGCAGA	97
10	(2) INFORMATION FOR SEQ ID NO:26:	
	(i) SEQUENCE CHARACTERISTICS:	
15	(A) LENGTH: 103 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
20	•	
	(ii) MOLECULE TYPE: DNA (genomic)	
	• •	
25		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	
	·	60
	CAGCCAGTAC ATGCTGAAGG TGTAGCCGCT GGCCTTGCAG CTCACCTTCA CGCTGGCGCC	60
	Chatchard Macroscot additions consorted eactasees	00
30	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG	103
30	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:	
	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:	
	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 105 base pairs	
	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 105 base pairs  (B) TYPE: nucleic acid	
35	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 105 base pairs	
35	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 105 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single	
35	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 105 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single	
35	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 105 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
35	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 105 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
35	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 105 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
35	GGGCTTCTTC ACCTCGGCGC CGCTCTGCAC CAGCTGCACC TGG  (2) INFORMATION FOR SEQ ID NO:27:  (i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 105 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	

5	CACCTTCAGC ATGTACTGGC TGGAGTGGGT GAAGCAGAGG CCCGGCCACG GCCTGGAGTG	60
	GGTGGGCGAG ATCAGCCCCG GCACCTTCAC CACCAACTAC AACGA	105
10	(2) INFORMATION FOR SEQ ID NO:28:	
	(i) SEQUENCE CHARACTERISTICS:	
15	(A) LENGTH: 107 base pairs	
	(B) TYPE: nucleic acid	,
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
20		
	(ii) MOLECULE TYPE: DNA (genomic)	
	•	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
	GTCCTCGCTG GTCAGGCTGC TCAGCTCCAT GTAGGCGGTG TTGGTGCTGG TGTCGGCGGT	60
30		
	GAAGGTGGCC TTGGCCTTGA ACTTCTCGTT GTAGTTGGTG GTGAAGG	107
	(2) INFORMATION FOR SEQ ID NO:29:	
35	·	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 83 base pairs	
	(B) TYPE: nucleic acid	
40	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
45		
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	
•		

•	AGCAGCCTGA CCAGCGAGGA CACCGCCGTG TACTACTGCG CCAGGTTCAG CCACTTCAGC	6.0
	GGCAGCAACT ACGACTACTT CGA	83
10	(2) INFORMATION FOR SEQ ID NO:30:	
	(i) SEQUENCE CHARACTERISTICS:	
15	(A) LENGTH: 83 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
20	(ii) MOLECULE TYPE: DNA (genomic)	
	·	
25		•
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
	TTTGGATCCT TCTAGAACTC ACCTGAGCTC ACGGTCACCA GGGTGCCCTG GCCCCAGTAG	60
30	TCGAAGTAGT CGTAGTTGCT GCC	83
35	(2) INFORMATION FOR SEQ ID NO:31:	
33	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 21 base pairs	
	(B) TYPE: nucleic acid	
40	(C) STRANDEDNESS: single	
•	(D) TOPOLOGY: linear	
45	(ii) MOLECULE TYPE: DNA (genomic)	
~		
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:	

5	TGAAGAAAGC TTGCCGCCAC C	21
	(2) INFORMATION FOR SEQ ID NO:32:	
10	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 23 base pairs	
	(B) TYPE: nucleic acid	
15	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
20		4
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:	
25		
	TTTGGATCCT TCTAGAACTC ACC	23
	(2) INFORMATION FOR SEQ ID NO:33:	
30		
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 24 base pairs	
25	(B) TYPE: nucleic acid	
35	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:	
45		
	TTTGGATCCT TCTAGAATAC TCAC	24
50	(2) INFORMATION FOR SEQ ID NO:34:	

	(i) SEQUENCE CHARACTERISTICS:	
5	(A) LENGTH: 16 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
10		
	(ii) MOLECULE TYPE: DNA (genomic)	
15		
75		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:	
20	AACAGCTATG ACCATG	16
	(2) INFORMATION FOR SEQ ID NO:35:	
25	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 30 base pairs	
	(B) TYPE: nucleic acid .	
	(C) STRANDEDNESS: single	
30	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
•		
35		•
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:	
40		
	CTGAACCTGT CGGGGATGCC GCTGATGCTC	30
	(2) INFORMATION FOR SEQ ID NO:36:	
45	(1) appropriate authoromorphisms	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 25 base pairs	
	(B) TYPE: nucleic acid	
50	(C) STRANDEDNESS: single	
	·	

5	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
10		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:	
		25
15	CCCGACAGGT TCAGCGGCAG CGGCA	23
	(2) INFORMATION FOR SEQ ID NO:37:	
20	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 22 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
25	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
30		
	·	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:	•
	•	
35	GGTCAGCACG ATCTCGCCGG TG	22
	(2) INFORMATION FOR SEQ ID NO:38:	
40		
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 27 base pairs	
	(B) TYPE: nucleic acid	
45	(C) STRANDEDNESS: single	·
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
50		
•		

5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:
	GAGATCGTGC TGACCCAGAG CCCCGGC
10	(2) INFORMATION FOR SEQ ID NO:39:
15	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 28 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear
20	(ii) MOLECULE TYPE: DNA (genomic)
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:
30	CCGGGGGCCT GCCTCACCCA CTCCAGCC  (2) INFORMATION FOR SEQ ID NO:40:
35	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 29 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>
45	(ii) MOLECULE TYPE: DNA (genomic)
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:

5	GAGGCAGGCC CCCGGCCACG GCCTGGAGT		29
	(2) INFORMATION FOR SEQ ID NO:41:		
10	(i) SEQUENCE CHARACTERISTICS:		
	(A) LENGTH: 34 base pairs		
	(B) TYPE: nucleic acid		
	(C) STRANDEDNESS: single		•
15	(D) TOPOLOGY: linear		
	(ii) MOLECULE TYPE: DNA (genomic)		·
20			
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:		
25	GAAGGTGGCC CTGGCCTTGA ACTTCTCGTT GTAG		34
	(2) INFORMATION FOR SEQ ID NO:42:	·	
30	(1) SEQUENCE CHARACTERISTICS:		4
	(A) LENGTH: 28 base pairs		
	(B) TYPE: nucleic acid		
	(C) STRANDEDNESS: single		
35	(D) TOPOLOGY: linear		
	(ii) MOLECULE TYPE: DNA (genomic)		
40			
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:		•
45	CAAGGCCAGG GCCACCTTCA CCGCCGAC		28
	(2) INFORMATION FOR SEQ ID NO:43:		
50			

	(1) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 31 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
10	(D) TOPOLOGY: linear	-
	(ii) MOLECULE TYPE: DNA (genomic)	
15		
	•	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:	
20	GTCCTCGCTC CTCAGGCTGC TCAGCTCCAT G	31
	didition eradocide renoticent o	3.
	(2) INFORMATION FOR SEQ ID NO:44:	
	(2) INFORMATION FOR SEQ ID NO. 44:	
25	(i) SEQUENCE CHARACTERISTICS:	
•	(A) LENGTH: 21 base pairs	
	(B) TYPE: nucleic acid	
30	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
35	•	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:	
40	•	
	CAGCCTGAGG AGCGAGGACA C	21
	(2) INFORMATION FOR SEQ ID NO:45:	
45		
	(i) SEQUENCE CHARACTERISTICS:	• •
	(A) LENGTH: 27 base pairs	
	(B) TYPE: nucleic acid	
50	(C) STRANDEDNESS: single	

. 60

		•	
5	(D) TOPOLOGY: linear	•	•
	(ii) MOLECULE TYPE: DNA (genomic)		
10			
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:		
15	CCATCCACTC CAGCCTCTGG CCGGGCC		27
	(2) INFORMATION FOR SEQ ID NO:46:	•	
20	(i) SEQUENCE CHARACTERISTICS:		
	(A) LENGTH: 33 base pairs		
	(B) TYPE: nucleic acid		
	(C) STRANDEDNESS: single		
25	(D) TOPOLOGY: linear		
:			
	(ii) MOLECULE TYPE: DNA (genomic)		
30			
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:		
35	CCATCCACTC CAGCCTCTGG CCGGGGGCCT GCC		33
	(2) INFORMATION FOR SEQ ID NO:47:		
40			
	(i) SEQUENCE CHARACTERISTICS:		
	(A) LENGTH: 27 base pairs		
	(B) TYPE: nucleic acid	·	
45	(C) STRANDEDNESS: single		
	(D) TOPOLOGY: linear		
	(ii) MOLECULE TYPE: DNA (genomic)	·	,
	(II) FORECODE IIIE. DIM (SCHOULE)		

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:	
10	CAGAGGCTGG AGTGGATGGG CGAGATC	27
	(2) INFÓRMATION FOR SEQ ID NO:48:	
15	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 20 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
20	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
25		•
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:	·
	GTGCTGGCGC TGGTGTCGGC	20
30		
	(2) INFORMATION FOR SEQ ID NO:49:	
	(i) SEQUENCE CHARACTERISTICS:	
35	(A) LENGTH: 21 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
40		
	(ii) MOLECULE TYPE: DNA (genomic)	
	•	
45	(x1) SEQUENCE DESCRIPTION: SEQ ID NO:49:	
	ACCAGCGCCA GCACCGCCTA C	21
	ACCAUCACE GUACACCIA C	21
50		

Claims

55

1. A reshaped human monoclonal antibody specific for IgE comprising an antigen binding site comprising,

in sequence, the hypervariable regions CDR1, CDR2 and CDR3; said CDR1 having the amino acid sequence Met-Tyr-Trp-Leu-Glu, said CDR2 having the amino acid sequence Glu-lle-Ser-Pro-Gly-Thr-Phe-Thr-Thr-Asn-Tyr-Asn-Glu-Lys-Phe-Lys-Ala, said CDR3 having the sequence Phe-Ser-His-Phe-Ser-Gly-Ser-Asn-Tyr-Asp-Tyr-Phe-Asp-Tyr-Phe-Asp-Tyr, said reshaped human antibody having an antigen binding affinity which at least about equals that of the murine CDR-donor antibody TES-C21, a direct equivalent or a derivative of said reshaped antibody.

- 2. A reshaped human antibody according to claim 1 comprising an antigen binding site comprising:
  - a) a first domain comprising, in sequence, the hypervariable regions CDR1, CDR2 and CDR3, said CDR1 having the amino acid sequence Met-Tyr-Trp-Leu-Glu, said CDR2 having the amino acid sequence Glu-lle-Ser-Pro-Gly-Thr-Phe-Thr-Thr-Asn-Tyr-Asn-Glu-Lys-Phe-Lys-Ala, said CDR3 having the amino acid sequence Phe-Ser-His-Phe-Ser-Gly-Ser-Asn-Tyr-Asp-Tyr-Phe-Asp-Tyr-Phe-Asp-Tyr, and
    - b) a second domain comprising, in sequence, the hypervariable regions CDR1, CDR2 and CDR3, said CDR1 having the amino acid sequence Arg-Ala-Ser-Gin-Ser-Ile-Giy-Thr-Asn-Ile-His, said CDR2 having the amino acid sequence Tyr-Ala-Ser-Giu-Ser, said CDR3 having the amino acid sequence Gin-Gin-Ser-Asp-Ser-Trp-Pro-Thr-Thr, said reshaped human antibody having an antigen binding affinity which at least about equals that of the murine CDR-donor antibody TES-C21, a direct equivalent or a derivative of said reshaped antibody.
- 3. A reshaped antibody according to claim 2 comprising:

10

15

25

30

45

- a) an immunoglobulin heavy chain or a fragment thereof which comprises a variable domain comprising in sequence the hypervariable regions CDR1_H, CDR2_H and CDR3_H and the constant part or fragment thereof of a human heavy chain, said CDR1_H having the amino acid sequence Met-Tyr-Trp-Leu-Glu, said CDR2_H having the amino acid sequence Glu-Ile-Ser-Pro-Gly-Thr-Phe-Thr-Thr-Asn-Tyr-Asn-Glu-Lys-Phe-Lys-Ala, said CDR3_H having the sequence Phe-Ser-His-Phe-Ser-Gly-Ser-Asn-Tyr-Asp-Tyr-Phe-Asp-Tyr-Phe-Asp-Tyr-Phe-Asp-Tyr-Rand
- b) an immunoglobulin light chain, or a fragment thereof, comprising a light chain variable domain comprising in sequence the hypervariable regions CDR1_L,CDR2_L and CDR3_L and the constant part, or a fragment thereof, of a human light chain, said CDR1_L having the amino acid sequence Arg-Ala-Ser-Gln-Ser-Ile-Gly-Thr-Asn-Ile-His, said CDR2_L having the amino acid sequence Tyr-Ala-Ser-Glu-Ser-Ile-Ser, said CDR3_L having the amino a sequence Gln-Gln-Ser-Asp-Ser-Trp-Pro-Thr-Thr, a direct equivalent or a derivative of said reshaped antibody.
- 4. A reshaped human antibody or a derivative thereof according to claim 2 comprising
  - a) a heavy chain comprising a variable domain having an amino acid sequence substantially identical with that shown in SEQ. ID. No. 11 starting with the amino acid at position 1 and ending with the amino acid at position 123 and the constant part of a human heavy chain; and
  - b) a light chain comprising a variable domain having an amino acid sequence substantially identical with that shown in SEQ. ID. No. 5 starting with the amino acid at position 1 and ending with the amino acid at position 107 and the constant part of a human light chain.
- 5. A reshaped human antibody or a derivative thereof according to claim 2 comprising:
  - a) a heavy chain comprising a variable domain having an amino acid sequence substantially identical with that shown in SEQ. ID. No. 13 starting with the amino acid at position 1 and ending with the amino acid at position 123 and the constant part of a human heavy chain; and
  - b) a light chain comprising a variable domain having an amino acid sequence substantially identical with that shown in SEQ. ID. No. 5 starting with the amino acid at position 1 and ending with the amino acid at position 107 and the constant part of a human light chain;
- A reshaped human antibody according to any of claims 1 to 5, wherein the heavy chain comprises the heavy chain human constant region γ1, and a derivative thereof.
  - A reshaped human antibody according to any of claims 3 to 5, wherein the heavy chain comprises the heavy chain human constant region γ1, and the light chain comprises the light chain human constant region κ, and derivatives thereof.
  - A reshaped human antibody according to claim 5 designated H3L1 produced by the cell line EH31.8, or a derivative thereof.

- A reshaped human antibody according to claim 4 designated H1L1 produced by the cell line EH11.13, or a derivative thereof
- 10. A derivative of a reshaped human antibody according to claim 1.
- 11. A process for the preparation of a reshaped human antibody, a direct equivalent or a derivative thereof according to claim 1 comprising culturing a suitable host producing a protein of the invention and, if required, isolating said protein and optionally converting it into a derivative.
- 12. A DNA construct encoding a heavy chain or a fragment thereof comprising
  a) a first part which encodes a variable domain comprising alternatively FRs and CDRs, said CDRs being in sequence CDR1_H, CDR2_H and CDR3_H, the amino acid sequences of which are identified in claim 3, and optionally,
  b) a second part encoding a human heavy chain constant part or fragment thereof.
- 13. A DNA construct encoding a light chain or a fragment thereof comprising:

  a) a first part which encodes a variable domain comprising alternatively FRs and CDRs, said CDRs being in sequence CDR1_L, CDR2_L and CDR3_L, the amino acid sequences of which are identified in claim 3, and optionally
  - b) a second part encoding a human light chain constant part or fragment thereof.
  - 14. A hybrid vector comprising a DNA construct according to claim 12 and/or a DNA construct according to claim 13.
  - 15. A host cell transformed with a hybrid vector according to claim 14.
  - 16. A host cell according to claim 15 which is cell line EH 11.13

20

25

- 17. A host cell according to claim 15 which is cell line EH 31.8
- 18. A reshaped human antibody, a direct equivalent thereof and/or a derivative thereof according to claim 1 for use in the prophylaxis and/or treatment of allergic reactions in humans.
  - 19. A pharmaceutical composition comprising an antibody, a direct equivalent and/or a derivative thereof according to claim 1, and a pharmaceutically acceptable carrier.
- 35 20. Use of an antibody, a direct equivalent and/or a derivative thereof according to claim 1 for the qualitative or quantitative determination of IgE.
  - 21. Test kit for the qualitative or quantitative determination of IgE comprising an antibody, a direct equivalent and/or a derivative thereof according to claim 1.

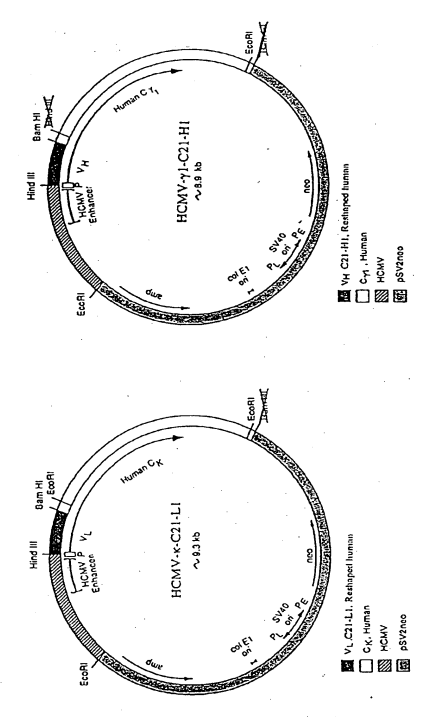


Fig. 1: HCMV-mammalian expression vectors



# **EUROPEAN SEARCH REPORT**

Application Number EP 93 81 0653

		ERED TO BE RELEVAN		
Category	Citation of document with ind of relevant pass		Relevant to claim	CLASSIFICATION OF THE APPLICATION (bcCLS)
D,Y	WO-A-89 06138 (TANOX * claims *	BIOSYSTEMS, INC.)	1-21	C12N15/13 C12P21/08 C12N5/10
Υ, σ	PROCEEDINGS OF THE N SCIENCES OF THE USA vol. 86, no. 24, De WASHINGTON DC, USA pages 10029 - 10033 C. QUEEN ET AL. 'A h binds to the interle * abstract *	cember 1989 , umanized antibody that	1-21	C12N15/62 A61K39/395 G01N33/577 G01N33/68
Y	EP-A-0 476 226 (SHIC KAISHA) * claims *	NOGI SEIYAKU KABUSHIKI	1-21	
Υ .	NATURE vol. 332 , 24 March pages 323 - 327 L. RIECHMANN ET AL. antibodies for thera * the whole document	'Reshaping human	1-21	TECHNICAL FIELDS SEARCHED (Int.CL.5)
A	INTERNATIONAL ARCHIVAPPLIED IMMUNOLOGY vol. 94, no. 1-4, 1 SWITZERLAND pages 87 - 90 C. HEUSSER ET AL. 'N regulation.' * abstract *	1991 , BASEL,	1-21	C12P A61K G01N C07K
		-/		
	The present search report has be	oce draws up for all claims		
<del></del>	Place of search	Date of campleties of the sourch		Domine
1	THE HAGUE	3 January 1994	No	oij, F
THE HAGUE  CATEGORY OF CITED DOCUMENTS  X: particularly relevant if taken alone Y: particularly relevant if combined with another document of the same category A: technological background O: non-written disclosure P: Lotermediate document		E : euriler patent éc after the filing (	in the application other reason	internet on, or



# **EUROPEAN SEARCH REPORT**

Application Number EP 93 81 0653

atogory	Citation of document with in- of relevant pas		Re to	event claim	CLASSIFICATION OF THE APPLICATION (IM.CLS)
<b>A</b>	GENE vol. 101, no. 2 , 19 NETHERLANDS pages 297 - 302 A. LEWIS ET AL. 'Ima complementarity-dete grafting by recombin	991 , AMSTERDAM, T munoglobulin ermining region		1	
	reaction to generate antibodies.' * the whole document	e humanised monocl	onal		
P,X	PROTEIN ENGINEERING vol. 6 SUPPL. , 1993 BRITAIN page 90		1-2	1	
	F. KOLBINGER ET AL. for the treatment of see the top abstract	fallergy.'	body		
P,Y	WO-A-92 17207 (TANO) * the whole document		) 1-2	1	TECHNICAL FIELDS SEARCHED (Int.Cl.5)
P,Y	WO-A-93 04173 (GENE * the whole document		1-2	1	
	·				
	The present search report has been drawn up for all claims				1
	Floor of search	Date of completion of the	- 1	N.	oij, F
THE HAGUE  CATEGORY OF CITED DOCUMENTS  X: particularly relevant if taken alone Y: particularly relevant if combined with another document of the same catagory A: technological background		after	1994 yor principle un or patent documen the filing date ment cited in the ment cited for eth	larlying C d, but pu	he invention hished on, or